to, a compound of the present invention in combination with: other (e.g., those other than inhibitors of the present invention) HIV protease inhibitors (e.g. Ro 31-8959, SC 52151, A-77003, A-80987, A-84538 and L-737,524); nucleoside and non-nucleoside reverse transcriptase inhibitors preferably nucleoside reverse transcriptase inhibitors such as AZT, DD1, d4T, DDC, 3TC, or PMEA, and non-nucleoside reverse transcriptase inhibitors such as nevirapine, pyridinones (e.g. L-697,661), BHAPs (e.g. U-90152), alpha-APA derivatives (e.g., R 18893) and TIBO derivatives (e.g. R82913); inhibitors of tat such as RO24-7429; drugs which inhibit binding of the virus to CD.sub.4 receptors; inhibitors of RNase, integrase, or rev; and immunomodulators such as IFN-.alpha. (.alpha.-interferon). DETD The antiviral activity of the retrocarbamate protease inhibitors of the present invention was evaluated by a microculture method which determines the increase in cell viability of an infected culture when a drug is added. The assay depends on the metabolic reduction of tetrazolium reagent by viable cells to yield a soluble colored formazan product. => d his (FILE 'HOME' ENTERED AT 08:25:40 ON 12 NOV 1997) FILE 'CAPLUS, EMBASE, MEDLINE, BIOSIS, USPATFULL' ENTERED AT 08:26:20 ON 12 NOV 1997 415 FILE CAPLUS 133 FILE EMBASE 141 FILE MEDLINE 251 FILE BIOSIS 10 FILE USPATFULL TOTAL FOR ALL FILES 950 S CYTOCHROME P450 MONOOXYGENASE? 0 FILE CAPLUS 0 FILE EMBASE O FILE MEDLINE L10 0 FILE BIOSIS O FILE USPATFULL L11 TOTAL FOR ALL FILES L12 0 S L6 AND RITONAVIR 74 FILE CAPLUS L13 159 FILE EMBASE L14L15 44 FILE MEDLINE L16 96 FILE BIOSIS 6 FILE USPATFULL L17 TOTAL FOR ALL FILES 379 S L6 AND SAQUINAVIR OR VX-478 OR MK-639 OR AG1343 OR A-77 L18 29 FILE CAPLUS L19 35 FILE EMBASE L20 40 FILE MEDLINE L21 31 FILE BIOSIS L22 4 FILE USPATFULL L23 TOTAL FOR ALL FILES

139 S L18 AND HIV PROTEASE INHIBITOR?

0 FILE CAPLUS

6 FILE EMBASE

L1

L2 L3

L4

L5

L6

L7

Г8

L9

L24

L25

L26

L27	19 FILE MEDLINE
L28	3 FILE BIOSIS
L29	2 FILE USPATFULL
	TOTAL FOR ALL FILES
L30	30 S L24 AND METABOLI?

nelfinavir, indinavir and VX-478. Clinically significant drug interactions have been predicted between ritonavir and a range of medications. In patients with HIV-1 infection, ritonavir markedly reduced viral load within 2 weeks of treatment onset and also increased CD4+ cell counts. In a large placebo-controlled trial in patients with advanced HIV infection, the addition of ritonavir to existing therapy reduced the risk of mortality by 43% and clinical progression by 56% after 6.1 months. Triple therapy with ritonavir plus zidovudine, in combination with lamivudine or zalcitabine, reduced HIV viraemia to below detectable levels in most patients with acute, and some patients with advanced HIV infection in 2 small trials. Early results suggest combination therapy with ritonavir and saquinavir increases CD4+ cell counts and decreases HIV RNA levels in patients with previously untreated HIV infection.

CTCheck Tags: Comparative Study; Human Administration, Oral *Anti-HIV Agents: PD, pharmacology Anti-HIV Agents: PK, pharmacokinetics Anti-HIV Agents: TU, therapeutic use Biological Availability Cytochrome P-450: GE, genetics Cytochrome P-450: ME, metabolism CD4-Positive T-Lymphocytes: CY, cytology *CD4-Positive T-Lymphocytes: DE, drug effects Dose-Response Relationship, Drug Double-Blind Method Drug Therapy, Combination Drug Tolerance *HIV Infections: DT, drug therapy HIV Infections: GE, genetics HIV Infections: MO, mortality *HIV Protease Inhibitors: PD, pharmacology HIV Protease Inhibitors: PK, pharmacokinetics HIV Protease Inhibitors: TU, therapeutic use Lamivudine: AD, administration & dosage Lamivudine: PD, pharmacology Lamivudine: TU, therapeutic use Leukocyte Count: DE, drug effects Randomized Controlled Trials Ritonavir: ME, metabolism *Ritonavir: PD, pharmacology Ritonavir: PK, pharmacokinetics Ritonavir: TU, therapeutic use RNA, Messenger: ME, metabolism Zalcitabine: AD, administration & dosage Zalcitabine: PD, pharmacology Zalcitabine: TU, therapeutic use Zidovudine: AD, administration & dosage Zidovudine: PD, pharmacology Zidovudine: TU, therapeutic use 0 (Anti-HIV Agents); 0 (HIV Protease Inhibitors); 0 (Ritonavir); 0 (RNA, Messenger)

L30 ANSWER 13 OF 30 MEDLINE

ACCESSION NUMBER: 96431786 MEDLINE

TITLE: Antiviral and resistance studies of AG1343, an orally bioavailable inhibitor of human

immunodeficiency virus protease.

AUTHOR: Patick A K; Mo H; Markowitz M; Appelt K; Wu B; Musick

L; Kalish V; Kaldor S; Reich S; Ho D; Webber S

CORPORATE SOURCE: Department of Pharmacology, Agouron Pharmaceuticals,

Inc., San Diego, CA 92121, USA.

SOURCE: ANTIMICROBIAL AGENTS AND CHEMOTHERAPY, (1996 Feb) 40

(2) 292-7.

Journal code: 6HK. ISSN: 0066-4804.

PUB. COUNTRY: United States

Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

FILE SEGMENT: Priority Journals

ENTRY MONTH: 9702 ENTRY WEEK: 19970204

AG1343 ([3S-(3R*, 4aR*, 8aR*, 2'S*, 3'S*)]-2-[2' hydroxy-3'-phenylthiomethyl-4'-aza-5'-oxo-5'-(2''-methyl-3''-hydro xy-phenyl) pentyl]-decahydroiso-quinoline-3-N-t-butylcarboxamide methanesulfonic acid) is a selective, nonpeptidic inhibitor of human immunodeficiency virus (HIV) protease (Ki = 2 nM) that was discovered by protein structure-based drug design methodologies. AG1343 was effective against the replication of several laboratory and clinical HIV type 1 (HIV-1) or HIV-2 isolates including pyridinone- and zidovudine-resistant strains, with 50% effective concentrations ranging from 9 to 60 nM. In reversibility studies, inhibition of gag (p55) proteolytic processing in HIV-1 particles from cells treated with AG1343 was maintained for up to 36 h after drug removal. The ability of virus to develop resistance to AG1343 was studied by serial passage of HIV-1 NL4.3 in the presence of increasing concentrations of drug. After 28 passages, a variant with a 30-fold reduction in susceptibility to AG1343 was isolated. Molecular analysis of the protease from this variant indicated a double change from a Met to Ile at residue 46 and an Ile to Val or Ala at residue 84 (M46I+I84V, A). Consistent with these findings, reductions in susceptibility were observed for recombinant viruses constructed to contain the single I84V change or the double M46I+I84V substitutions. Resistance, however, was not detected for recombinant viruses containing other key mutations in HIV-1 protease, including a Val to Ile change at residue 32 or a Val to Ala or Phe at residue

TI Antiviral and resistance studies of AG1343, an orally bioavailable inhibitor of human immunodeficiency virus protease.

82. The potent anti-HIV activity of **AG1343** against several isolates suggests that **AG1343** should perform well during

ongoing human phase II clinical trials.

AG1343 ([3S-(3R*, 4aR*, 8aR*, 2'S*, 3'S*)]-2-[2' hydroxy-3'-phenylthiomethyl-4'-aza-5'-oxo-5'-(2''-methyl-3''-hydro xy-phenyl) pentyl]-decahydroiso-quinoline-3-N-t-butylcarboxamide methanesulfonic acid) is a selective, nonpeptidic inhibitor of human immunodeficiency virus (HIV) protease (Ki = 2 nM) that was discovered by protein structure-based drug design methodologies.

AG1343 was effective against the replication of several laboratory and clinical HIV type 1 (HIV-1) or HIV-2 isolates including pyridinone- and zidovudine-resistant strains, with 50% effective concentrations ranging from 9 to 60 nM. In reversibility studies, inhibition of gag (p55) proteolytic processing in HIV-1 particles from cells treated with AG1343 was maintained for up to 36 h after drug removal. The ability of virus to develop resistance to AG1343 was studied by serial passage of

HIV-1 NL4.3 in the presence of increasing concentrations of drug. After 28 passages, a variant with a 30-fold reduction in susceptibility to AG1343 was isolated. Molecular analysis of the protease from this variant indicated a double change from a Met to Ile at residue 46 and an Ile to Val or Ala at residue 84 (M46I+I84V, A). Consistent with these findings, reductions in susceptibility were observed for recombinant viruses constructed to contain the single I84V change or the double M46I+I84V substitutions. Resistance, however, was not detected for recombinant viruses containing other key mutations in HIV-1 protease, including a Val to Ile change at residue 32 or a Val to Ala or Phe at residue 82. The potent anti-HIV activity of AG1343 against several isolates suggests that AG1343 should perform well during ongoing human phase II clinical trials. CTCheck Tags: Comparative Study Amino Acid Sequence *Antiviral Agents: PD, pharmacology Cells, Cultured Drug Resistance, Microbial Gene Products, gag: ME, metabolism *HIV Protease Inhibitors: PD, pharmacology *HIV-1: DE, drug effects HIV-1: EN, enzymology HIV-1: GE, genetics *HIV-2: DE, drug effects *Isoquinolines: PD, pharmacology Microbial Sensitivity Tests Molecular Sequence Data Reverse Transcriptase Inhibitors: PD, pharmacology Saquinavir: PD, pharmacology *Sulfonic Acids: PD, pharmacology Zidovudine: PD, pharmacology 0 (Antiviral Agents); 0 (AG 1343); 0 (Gene Products, gag); 0 (CN HIV Protease Inhibitors); 0 (Isoquinolines); 0 (Reverse Transcriptase Inhibitors); 0 (Sulfonic Acids) L30 ANSWER 14 OF 30 MEDLINE ACCESSION NUMBER: 96417630 MEDLINE TITLE: Role of cytochrome P450 3A4 in human metabolism of MK-639, a potent human immunodeficiency virus protease inhibitor. Chiba M; Hensleigh M; Nishime J A; Balani S K; Lin J AUTHOR: CORPORATE SOURCE: Department of Drug Metabolism, Merck Research Laboratories, West Point, PA 19486, USA. DRUG METABOLISM AND DISPOSITION, (1996 Mar) 24 (3) SOURCE: 307-14. Journal code: EBR. ISSN: 0090-9556. PUB. COUNTRY: United States Journal; Article; (JOURNAL ARTICLE) LANGUAGE: English Priority Journals FILE SEGMENT: ENTRY MONTH: 9702 ENTRY WEEK: 19970204 MK-639 (L-735,524) is a potent human immunodeficiency virus protease inhibitor under investigation in the

treatment of acquired immunodeficiency syndrome. Five in vitro approaches have been used to identify the cytochrome P450 isoform(s) responsible for the human microsomal oxidative metabolism of MK-639. These approaches are: 1) chemical inhibition; 2) immunochemical inhibition; 3) metabolism by cDNA-expressed human cytochrome P450 enzymes; 4) a correlation analysis; and 5) competitive inhibition of marker activities. Ketoconazole and troleandomycin, both selective inhibitors for cytochrome P450 3A4 (CYP3A4), markedly inhibited the formation of all oxidative metabolites of MK-639; whereas other inhibitors (furafylline, sulfaphenazole, quinidine, S-mephenytoin, and diethyldithiocarbamate) had little effect on MK-639 metabolism. This suggested the involvement of CYP3A4 in MK-639 metabolism. Consistent with this, an anti-rat CYP3A1 rabbit polyclonal antibody, which shows a cross-reactive inhibition of CYP3A4-dependent testosterone 6beta-hydroxylation in human liver microsomes, completely inhibited MK-639 metabolism. Human recombinant CYP3A4 showed a high metabolic activity to form all MK-639 metabolites found in native human liver microsomes. In addition, the formation of individual MK-639 metabolites correlated well with each other and with testosterone 6beta-hydroxylation in 12 different human liver microsomes, whereas no correlation was observed between MK -639 metabolite formation and bufuralol 1'-hydroxylation (or tolbutamide methyl hydroxylation). Furthermore, MK-639 strongly inhibited testosterone 6beta-hydroxylation in a concentration-dependent manner. Kinetic analysis showed that MK-639 is a very potent competitive inhibitor for testosterone 6beta-hydroxylation, with a Ki value of approximately 0.5 mu M. Collectively, these results consistently indicate that CYP3A4 is the isoform responsible for the oxidative metabolism of MK-639 in human liver microsomes. Role of cytochrome P450 3A4 in human metabolism of

- TI Role of cytochrome P450 3A4 in human metabolism of MK-639, a potent human immunodeficiency virus protease inhibitor.
- MK-639 (L-735,524) is a potent human AB immunodeficiency virus protease inhibitor under investigation in the treatment of acquired immunodeficiency syndrome. Five in vitro approaches have been used to identify the cytochrome P450 isoform(s) responsible for the human microsomal oxidative metabolism of MK-639. These approaches are: 1) chemical inhibition; 2) immunochemical inhibition; 3) metabolism by cDNA-expressed human cytochrome P450 enzymes; 4) a correlation analysis; and 5) competitive inhibition of marker activities. Ketoconazole and troleandomycin, both selective inhibitors for cytochrome P450 3A4 (CYP3A4), markedly inhibited the formation of all oxidative metabolites of MK-639; whereas other inhibitors (furafylline, sulfaphenazole, quinidine, S-mephenytoin, and diethyldithiocarbamate) had little effect on MK-639 metabolism. This suggested the involvement of CYP3A4 in MK-639 metabolism. Consistent with this, an anti-rat CYP3A1 rabbit polyclonal antibody, which shows a cross-reactive inhibition of CYP3A4-dependent testosterone 6beta-hydroxylation in human liver microsomes, completely inhibited MK-639

metabolism. Human recombinant CYP3A4 showed a high metabolic activity to form all MK-639 metabolites found in native human liver microsomes. In addition, the formation of individual MK-639 metabolites correlated well with each other and with testosterone 6beta-hydroxylation in 12 different human liver microsomes, whereas no correlation was observed between MK -639 metabolite formation and bufuralol 1'-hydroxylation (or tolbutamide methyl hydroxylation). Furthermore, MK-639 strongly inhibited testosterone 6beta-hydroxylation in a concentration-dependent manner. Kinetic analysis showed that MK-639 is a very potent competitive inhibitor for testosterone 6beta-hydroxylation, with a Ki value of approximately 0.5 mu M. Collectively, these results consistently indicate that CYP3A4 is the isoform responsible for the oxidative metabolism of MK-639 in human liver microsomes. Check Tags: Human Antibiotics, Macrolide: PD, pharmacology Antifungal Agents: PD, pharmacology *Cytochrome P-450: ME, metabolism *Hydroxylases: ME, metabolism Hydroxylation: DE, drug effects *HIV Protease Inhibitors: ME, metabolism *Indinavir: ME, metabolism Isoenzymes Ketoconazole: PD, pharmacology *Microsomes, Liver: ME, metabolism Troleandomycin: PD, pharmacology EC 1.14. (Hydroxylases); EC 1.14.99.- (nifedipine oxidase); 0 (Antibiotics, Macrolide); 0 (Antifungal Agents); 0 (HIV Protease Inhibitors); 0 (Isoenzymes) L30 ANSWER 15 OF 30 MEDLINE ACCESSION NUMBER: 96338373 MEDLINE TITLE: Human serum alpha 1 acid glycoprotein reduces uptake, intracellular concentration, and antiviral activity of A-80987, an inhibitor of the human immunodeficiency virus type 1 protease. Bilello J A; Bilello P A; Stellrecht K; Leonard J; AUTHOR: Norbeck D W; Kempf D J; Robins T; Drusano G L Department of Medicine, Albany Medical College, New CORPORATE SOURCE: York 12208, USA. ANTIMICROBIAL AGENTS AND CHEMOTHERAPY, (1996 Jun) 40 SOURCE: (6) 1491-7. Journal code: 6HK. ISSN: 0066-4804. PUB. COUNTRY: United States Journal; Article; (JOURNAL ARTICLE) LANGUAGE: English Priority Journals FILE SEGMENT: ENTRY MONTH: 9704

CT

ENTRY WEEK: 19970401 The therapeutic utility of a human immunodeficiency virus type 1 (HIV-1) protease inhibitor may depend on its intracellular concentration, which is a property of its uptake, metabolism , and/or efflux. Previous studies in our laboratory indicated that the addition of alpha 1 acid glycoprotein (alpha 1 AGP) to the medium markedly increased the amount of the drug required to limit

infection in vitro. In this study, physiologically relevant concentrations of alpha 1 AGP and a radiolabeled inhibitor, A-80987, were used to determine both the uptake and activity of the agent in HIV-1-infected human peripheral blood mononuclear cells and cell lines. Both the uptake and efflux of 14C-labeled A-80987 were rapid (t1/2, < 5 min). Uptake of the drug was linearly dependent on the concentration but insensitive to the metabolic inhibitors KF, sodium cyanide, or CCCP (carbonyl cyanide m-chlorophenyl hydrazone). The amount of A-80987 which entered the cells was inversely proportional to the concentration of alpha 1 AGP (r2, 0.99) and directly proportional to the amount of extracellular non-protein-bound drug (r2, 0.99). Most importantly, the antiviral activity of the drug in HIV-1-infected peripheral blood mononuclear cells and MT-2 cells was directly related to the amount of intracellular A-80987. This study demonstrates that A-80987 binds to alpha 1 AGP, resulting in a free fraction below 10%. Cellular uptake of A-80987 is proportionally decreased in the presence of alpha 1 AGP, which results in less-than-expected antiviral activity. Importantly, we demonstrate for the first time that the inhibition of HIV protease is highly correlated with the amount of intracellular inhibitor.

- TI Human serum alpha 1 acid glycoprotein reduces uptake, intracellular concentration, and antiviral activity of A-80987
- , an inhibitor of the human immunodeficiency virus type 1 protease. The therapeutic utility of a human immunodeficiency virus type 1 AB (HIV-1) protease inhibitor may depend on its intracellular concentration, which is a property of its uptake, metabolism , and/or efflux. Previous studies in our laboratory indicated that the addition of alpha 1 acid glycoprotein (alpha 1 AGP) to the medium markedly increased the amount of the drug required to limit infection in vitro. In this study, physiologically relevant concentrations of alpha 1 AGP and a radiolabeled inhibitor, A-80987, were used to determine both the uptake and activity of the agent in HIV-1-infected human peripheral blood mononuclear cells and cell lines. Both the uptake and efflux of 14C-labeled A-80987 were rapid (t1/2, < 5 min). Uptake of the drug was linearly dependent on the concentration but insensitive to the metabolic inhibitors KF, sodium

amount of A-80987 which entered the cells was inversely proportional to the concentration of alpha 1 AGP (r2, 0.99) and directly proportional to the amount of extracellular non-protein-bound drug (r2, 0.99). Most importantly, the antiviral activity of the drug in HIV-1-infected peripheral blood mononuclear cells and MT-2 cells was directly related to the amount of intracellular A-80987. This study demonstrates

cyanide, or CCCP (carbonyl cyanide m-chlorophenyl hydrazone). The

that A-80987 binds to alpha 1 AGP, resulting in a free fraction below 10%. Cellular uptake of A-

80987 is proportionally decreased in the presence of alpha 1 AGP, which results in less-than-expected antiviral activity. Importantly, we demonstrate for the first time that the inhibition of HIV protease is highly correlated with the amount of intracellular inhibitor.

CT Check Tags: Human

Cell Line

HIV Protease Inhibitors: ME, metabolism

*HIV Protease Inhibitors: PK, pharmacokinetics *HIV-1: DE, drug effects HIV-1: ME, metabolism Orosomucoid: ME, metabolism *Orosomucoid: PD, pharmacology Polymerase Chain Reaction Protein Binding Pyridines: ME, metabolism *Pyridines: PK, pharmacokinetics RNA, Viral: DE, drug effects 0 (A 80987); 0 (HIV Protease Inhibitors); 0 (Orosomucoid); 0 (Pyridines); 0 (RNA, Viral) L30 ANSWER 16 OF 30 MEDLINE ACCESSION NUMBER: 96298219 MEDLINE TITLE: In vitro metabolism of a potent HIV -protease inhibitor (141W94) using rat, monkey and human liver S9. Singh R; Chang S Y; Taylor L C AUTHOR: CORPORATE SOURCE: Glaxo Wellcome Inc., Research Triangle Park, NC 27709, USA. RAPID COMMUNICATIONS IN MASS SPECTROMETRY, (1996) 10 SOURCE: (9) 1019-26. Journal code: A9Q. ISSN: 0951-4198. PUB. COUNTRY: ENGLAND: United Kingdom Journal; Article; (JOURNAL ARTICLE) LANGUAGE: English FILE SEGMENT: Priority Journals ENTRY MONTH: 9704 ENTRY WEEK: 19970403 Compound 141W94 (Vertex VX478) (3S)-tetrahydro-3-furyl N-[((S,2R)-3-(4-amino-N-isobutylbenzenesulfonamido)-1-benzyl-2-hydroxypropyl] carbamate, is a potent HIVprotease inhibitor and is currently undergoing clinical trials. The purpose of this study was the rapid identification of the phase I and II in vitro metabolite of 141W94 using mass spectrometry. Four different sources of liver S9 fractions were used for studying comparative in vitro metabolism of 141W94. They were obtained from Arochlor-induced rat, normal (untreated) rat, cynomolgus monkey and human livers. Selected incubations were supplemented with uridine diphosphate glucuronic acid and the reduced form of glutathione. The predominant species seen in the incubation mixture was the parent compound 141W94. Metabolites arising from ring opening to form the diol and carboxylic acid and oxidation of the tetrahydrofurran ring (formation of dihydrofuran) were identified. In addition, of the two monohydroxylated products identified, one resulted from hydroxylation on the aniline ring and the other from hydroxylation at the benzylic position. Two different glucuronides were also observed. Comparing the three species, very little metabolism was seen in the normal (non-induced) rat. The metabolic profile and extent of metabolism with induced rat, monkey and human S9 was similar. Induced rat S9 incubation showed the formation of two unique metabolites that were not seen in non-induced rat, monkey and human S9 fractions. They were the monohydroxylated glucuronide and a carbamate cleavage product. The metabolites were

identified using mass spectrometry based on their molecular masses

and fragmentation patterns. ΤI In vitro metabolism of a potent HIVprotease inhibitor (141W94) using rat, monkey and human liver S9. AB Compound 141W94 (Vertex VX478) (3S)-tetrahydro-3-furyl N-[((S,2R)-3-(4-amino-N-isobutylbenzenesulfonamido)-1-benzyl-2-hydroxypropyl] carbamate, is a potent HIVprotease inhibitor and is currently undergoing clinical trials. The purpose of this study was the rapid identification of the phase I and II in vitro metabolite of 141W94 using mass spectrometry. Four different sources of liver S9 fractions were used for studying comparative in vitro metabolism of 141W94. They were obtained from Arochlor-induced rat, normal (untreated) rat, cynomolgus monkey and human livers. Selected incubations were supplemented with uridine diphosphate glucuronic acid and the reduced form of glutathione. The predominant species seen in the incubation mixture was the parent compound 141W94. Metabolites arising from ring opening to form the diol and carboxylic acid and oxidation of the tetrahydrofurran ring (formation of dihydrofuran) were identified. In addition, of the two monohydroxylated products identified, one resulted from hydroxylation on the aniline ring and the other from hydroxylation at the benzylic position. Two different glucuronides were also observed. Comparing the three species, very little metabolism was seen in the normal (non-induced) rat. The metabolic profile and extent of metabolism with induced rat, monkey and human S9 was similar. Induced rat S9 incubation showed the formation of two unique metabolites that were not seen in non-induced rat, monkey and human S9 fractions. They were the monohydroxylated glucuronide and a carbamate cleavage product. The metabolites were identified using mass spectrometry based on their molecular masses and fragmentation patterns. CTCheck Tags: Animal; Comparative Study; Human; In Vitro Aroclors: PD, pharmacology Chromatography, High Pressure Liquid *HIV Protease Inhibitors: ME, metabolism Liver: CY, cytology *Liver: ME, metabolism Macaca fascicularis Rats Rats, Sprague-Dawley Species Specificity Spectrophotometry, Ultraviolet Spectrum Analysis, Mass Subcellular Fractions: ME, metabolism *Sulfonamides: ME, metabolism CN 0 (Aroclors); 0 (HIV Protease Inhibitors); 0 (Sulfonamides); 0 (**VX 478**) L30 ANSWER 17 OF 30 MEDLINE ACCESSION NUMBER: 96279343 MEDLINE TITLE: Kinetic characterization of human immunodeficiency

Rao B G; Livingston D J

AUTHOR:

CORPORATE SOURCE: Vertex Pharmaceuticals Incorporated, Cambridge,

virus type-1 protease-resistant variants.

Pazhanisamy S; Stuver C M; Cullinan A B; Margolin N;

Massachusetts 02139, USA.

SOURCE: JOURNAL OF BIOLOGICAL CHEMISTRY, (1996 Jul 26) 271

(30) 17979-85.

Journal code: HIV. ISSN: 0021-9258.

PUB. COUNTRY:

United States

Journal; Article; (JOURNAL ARTICLE)

LANGUAGE:

English

FILE SEGMENT:

Priority Journals; Cancer Journals

ENTRY MONTH: 9611

Passage of human immunodeficiency virus type-1 (HIV-1) in T-lymphocyte cell lines in the presence of increasing concentrations of the hydroxylethylamino sulfonamide inhibitor vx-478 or VB-11328 results in sequential accumulation of mutations in HIV-1 protease. We have characterized recombinant HIV-1 proteases that contain these mutations either individually (L10F, M46I, I47V, I50V) or in combination (the double mutant L10F/I50V and the triple mutant M46I/I47V/I50V). The catalytic properties and affinities for sulfonamide inhibitors and other classes of inhibitors were determined. For the I50V mutant, the efficiency (kcat/Km) of processing peptides designed to mimic cleavage junctions in the HIV-1 gag-pol polypeptide was decreased up to 25-fold. The triple mutant had a 2-fold higher processing efficiency than the I50V single mutant for peptide substrates with Phe/Pro and Tyr/Pro cleavage sites, suggesting that the M46I and I47V mutations are compensatory. The effects of mutation on processing efficiency were used in conjunction with the inhibition constant (Ki) to evaluate the advantage of the mutation for viral replication in the presence of drug. These analyses support the virological observation that the addition of M46I and I47V mutations on the I50V mutant background enables increased survival of the HIV-1 virus as it replicates in the presence of VX-478. Crystal structures and molecular models of the active site of the HIV-1 protease mutants suggest that changes in the active site can selectively affect the binding energy of inhibitors with little corresponding change in substrate binding.

AB Passage of human immunodeficiency virus type-1 (HIV-1) in T-lymphocyte cell lines in the presence of increasing concentrations of the hydroxylethylamino sulfonamide inhibitor vx-478 or VB-11328 results in sequential accumulation of mutations in HIV-1 protease. We have characterized recombinant HIV-1 proteases that contain these mutations either individually (L10F, M46I, I47V, I50V) or in combination (the double mutant L10F/I50V and the triple mutant M46I/I47V/I50V). The catalytic properties and affinities for sulfonamide inhibitors and other classes of inhibitors were determined. For the I50V mutant, the efficiency (kcat/Km) of processing peptides designed to mimic cleavage junctions in the HIV-1 gag-pol polypeptide was decreased up to 25-fold. The triple mutant had a 2-fold higher processing efficiency than the I50V single mutant for peptide substrates with Phe/Pro and Tyr/Pro cleavage sites, suggesting that the M46I and I47V mutations are compensatory. The effects of mutation on processing efficiency were used in conjunction with the inhibition constant (Ki) to evaluate the advantage of the mutation for viral replication in the presence of drug. These analyses support the virological observation that the addition of M46I and I47V mutations on the I50V mutant background enables increased survival of the HIV-1 virus as it replicates in the presence of VX-478. Crystal structures and molecular models of the active site of the HIV-1 protease mutants suggest that changes in the active site can

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selectively affect the binding energy of inhibitors with little
     corresponding change in substrate binding.
CT
     Check Tags: Comparative Study
      Amino Acid Sequence
      Binding Sites
      Hydrolysis
      HIV Protease: DE, drug effects
     *HIV Protease: GE, genetics
      HIV Protease Inhibitors: PD, pharmacology
     *HIV-1: EN, enzymology
     *HIV-1: GE, genetics
      Isoquinolines: PD, pharmacology
      Kinetics
      Models, Molecular
      Molecular Sequence Data
     *Mutation
      Oligopeptides: ME, metabolism
      Pyridines: PD, pharmacology
      Quinolines: PD, pharmacology
      Selection (Genetics)
      Substrate Specificity
      Sulfonamides: PD, pharmacology
      Variation (Genetics)
     EC 3.4.23.- (HIV Protease); 0 (HIV Protease
CN
     Inhibitors); 0 (Isoquinolines); 0 (Oligopeptides); 0
     (Pyridines); 0 (Quinolines); 0 (Sulfonamides); 0 (VX
     478)
L30 ANSWER 18 OF 30 MEDLINE
ACCESSION NUMBER:
                    96217762
                                 MEDLINE
TITLE:
                    Relevance of plasma protein binding to antiviral
                    activity and clinical efficacy of inhibitors of human
                    immunodeficiency virus protease [letter; comment].
COMMENT:
                    Comment on: J Infect Dis 1995 Nov; 172(5):1238-45
                    Bilello J A; Drusano G L
AUTHOR:
SOURCE:
                    JOURNAL OF INFECTIOUS DISEASES, (1996 Jun) 173 (6)
                    1524-6.
                    Journal code: IH3. ISSN: 0022-1899.
PUB. COUNTRY:
                    United States
                    Commentary
                    Letter
LANGUAGE:
                    English
FILE SEGMENT:
                    Abridged Index Medicus Journals; Priority Journals
ENTRY MONTH:
                    9609
CT
    Check Tags: Human
     Antiviral Agents: ME, metabolism
     *Antiviral Agents: PD, pharmacology
     *Blood Proteins: ME, metabolism
      Clinical Trials
     *HIV: DE, drug effects
     HIV Infections: DT, drug therapy
     HIV Protease Inhibitors: ME, metabolism
     *HIV Protease Inhibitors: PD, pharmacology
     Orosomucoid: ME, metabolism
      Protein Binding
     Sulfonamides: ME, metabolism
     *Sulfonamides: PD, pharmacology
     0 (Antiviral Agents); 0 (Blood Proteins); 0 (HIV
CN
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Protease Inhibitors); 0 (Orosomucoid); 0 (Sulfonamides); 0 (VX 478)

L30 ANSWER 19 OF 30 MEDLINE

ACCESSION NUMBER: 96202332 MEDLINE

TITLE: Human immunodeficiency virus type 1 viral background

plays a major role in development of resistance to

protease inhibitors.

AUTHOR: Rose R E; Gong Y F; Greytok J A; Bechtold C M; Terry

B J; Robinson B S; Alam M; Colonno R J; Lin P F

CORPORATE SOURCE: Department of Virology, Bristol-Myers Squibb Company,

Wallingford, CT 06492, USA.

PROCEEDINGS OF THE NATIONAL ACADEMY OF SCIENCES OF SOURCE:

THE UNITED STATES OF AMERICA, (1996 Feb 20) 93 (4)

1648-53.

Journal code: PV3. ISSN: 0027-8424.

PUB. COUNTRY: United States

Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

FILE SEGMENT: Priority Journals; Cancer Journals

ENTRY MONTH: 9609

The observed in vitro and in vivo benefit of combination treatment with anti-human immunodeficiency virus (HIV) agents prompted us to examine the potential of resistance development when two protease inhibitors are used concurrently. Recombinant HIV-1 (NL4-3) proteases containing combined resistance mutations associated with BMS-186318 and A-77003 (or saquinavir) were either inactive or had impaired enzyme activity. Subsequent construction of HIV-1 (NL4-3) proviral clones containing the same mutations yielded viruses that were severely impaired in growth or nonviable, confirming that combination therapy may be advantageous. However, passage of BMS-186318-resistant HIV-1 (RF) in the presence of either saquinavir or SC52151, which represented sequential drug treatment, produced viable viruses resistant to both BMS-186318 and the second compound. The predominant breakthrough virus contained the G48V/A71T/V82A protease mutations. The clone-purified RF (G48V/A71T/V82A) virus, unlike the corresponding defective NL4-3 triple mutant, grew well and displayed cross-resistance to four distinct protease inhibitors. Chimeric virus and in vitro mutagenesis studies indicated that the RF-specific protease sequence, specifically the Ile at residue 10, enabled the NL4-3 strain with the triple mutant to grow. Our results clearly indicate that viral genetic background will play a key role in determining whether cross-resistance variants will arise.

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SOURCE:

the G48V/A71T/V82A protease mutations. The clone-purified RF (G48V/A71T/V82A) virus, unlike the corresponding defective NL4-3 triple mutant, grew well and displayed cross-resistance to four distinct protease inhibitors. Chimeric virus and in vitro mutagenesis studies indicated that the RF-specific protease sequence, specifically the Ile at residue 10, enabled the NL4-3 strain with the triple mutant to grow. Our results clearly indicate that viral genetic background will play a key role in determining whether cross-resistance variants will arise. Check Tags: Comparative Study; Human Amino Acid Sequence Carbamates: PD, pharmacology Clone Cells Drug Administration Schedule Drug Resistance, Microbial: GE, genetics Drug Therapy, Combination DNA Mutational Analysis DNA, Recombinant: GE, genetics DNA, Viral: GE, genetics Ethanolamines: PD, pharmacology Hela Cells *HIV Protease: GE, genetics HIV Protease Inhibitors: AD, administration & dosage *HIV Protease Inhibitors: PD, pharmacology *HIV-1: DE, drug effects HIV-1: EN, enzymology HIV-1: GE, genetics Isoquinolines: PD, pharmacology Methylurea Compounds: PD, pharmacology Molecular Sequence Data Point Mutation Proviruses: EN, enzymology Proviruses: GE, genetics Pyridines: PD, pharmacology Quinolines: PD, pharmacology Recombinant Fusion Proteins: AI, antagonists & inhibitors Recombinant Fusion Proteins: ME, metabolism T-Lymphocytes Urea: AA, analogs & derivatives Urea: PD, pharmacology EC 3.4.23.- (HIV Protease); 0 (BMS 186318); 0 (Carbamates); 0 (DNA, Recombinant); 0 (DNA, Viral); 0 (Ethanolamines); 0 (HIV Protease Inhibitors); 0 (Isoquinolines); 0 (Methylurea Compounds); 0 (Pyridines); 0 (Quinolines); 0 (Recombinant Fusion Proteins); 0 (SC 52151) L30 ANSWER 20 OF 30 MEDLINE ACCESSION NUMBER: 96139551 MEDLINE Efficacy of constant infusion of A-TITLE: 77003, an inhibitor of the human immunodeficiency virus type 1 (HIV-1) protease, in limiting acute HIV-1 infection in vitro. Bilello J A; Bilello P A; Kort J J; Dudley M N; AUTHOR: Leonard J; Drusano G L CORPORATE SOURCE: Department of Medicine, Albany Medical College, New York 12208, USA.

ANTIMICROBIAL AGENTS AND CHEMOTHERAPY, (1995 Nov) 39

(11) 2523-7.

Journal code: 6HK. ISSN: 0066-4804.

PUB. COUNTRY: United States

Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

FILE SEGMENT: Priority Journals

ENTRY MONTH: 9605

A-77003, a human immunodeficiency virus type 1 (HIV-1) protease inhibitor, is effective for both acute and chronic infection in vitro and was evaluated clinically by continuous intravenous infusion administration. The minimum effective dose (the concentration required to completely inhibit viral replication) was determined in vitro in a population of uninfected (99%) and HIV-infected (1%) cells exposed to A-77003 by continuous infusion in hollow-fiber bioreactors. The production of infectious HIV and release of p24 antigen from infected cells were completely inhibited in cultures exposed to A-77003 at or above a concentration of 0.5 microM. Measurement of unintegrated HIV-1 DNA synthesis and flow cytometric analysis for cells expressing HIV p24 antigen demonstrated that the spread of HIV to uninfected cells was also blocked at 0.5 microM A-77003. Dose deescalation to 0.25 microM or removal of A-77003 resulted in the limited spread of the virus throughout the culture, the resumption of viral DNA synthesis, and release of p24. HIV produced after exposure to 0.5 microM A-77003 was noninfectious for a period of 72 h after the removal of the drug. Addition of 1 mg of alpha 1-acid glycoprotein per ml to this in vitro system completely ablated the anti-HIV effect of 0.5 microM A-77003. These data suggest that determination of the minimum effective dose under conditions which simulate human pharmacodynamic patterns may be useful in determining the initial dose and schedule for clinical trials. However, other factors, such as serum protein binding, may influence the selection of a therapeutic regimen.

TI Efficacy of constant infusion of A-77003, an inhibitor of the human immunodeficiency virus type 1 (HIV-1) protease, in limiting acute HIV-1 infection in vitro.

AB A-77003, a human immunodeficiency virus type 1 (HIV-1) protease inhibitor, is effective for both acute and chronic infection in vitro and was evaluated clinically by continuous intravenous infusion administration. The minimum effective dose (the concentration required to completely inhibit viral replication) was determined in vitro in a population of uninfected (99%) and HIV-infected (1%) cells exposed to A-77003 by continuous infusion in hollow-fiber bioreactors. The production of infectious HIV and release of p24 antigen from infected cells were completely inhibited in cultures exposed to A-77003 at or above a concentration of 0.5 microM. Measurement of unintegrated HIV-1 DNA synthesis and flow cytometric analysis for cells expressing HIV p24 antigen demonstrated that the spread of HIV to uninfected cells was also blocked at 0.5 microM A-77003. Dose deescalation to 0.25 microM or removal of A-77003 resulted in the limited spread of the virus throughout the culture, the resumption of viral DNA synthesis, and release of p24. HIV produced after exposure to 0.5 microM A-77003 was noninfectious for a period of 72 h after the removal of the drug. Addition of 1 mg of alpha 1-acid glycoprotein per ml to this in vitro system completely ablated the anti-HIV effect of 0.5 microM A-77003. These

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conditions which simulate human pharmacodynamic patterns may be useful in determining the initial dose and schedule for clinical trials. However, other factors, such as serum protein binding, may influence the selection of a therapeutic regimen. CTCheck Tags: Human Antiviral Agents: AD, administration & dosage *Antiviral Agents: PD, pharmacology Cell Line Dose-Response Relationship, Drug DNA, Viral: BI, biosynthesis Flow Cytometry HIV Core Protein p24: ME, metabolism HIV Protease Inhibitors: AD, administration & dosage *HIV Protease Inhibitors: PD, pharmacology *HIV-1: DE, drug effects HIV-1: PH, physiology Methylurea Compounds: AD, administration & dosage *Methylurea Compounds: PD, pharmacology Orosomucoid: ME, metabolism Orosomucoid: PD, pharmacology Polymerase Chain Reaction Pyridines: AD, administration & dosage *Pyridines: PD, pharmacology T-Lymphocytes: VI, virology Virus Replication: DE, drug effects CN 0 (Antiviral Agents); 0 (DNA, Viral); 0 (HIV Core Protein p24); 0 (HIV Protease Inhibitors); 0 (Methylurea Compounds); 0 (Orosomucoid); 0 (Pyridines) L30 ANSWER 21 OF 30 MEDLINE ACCESSION NUMBER: 96036379 MEDLINE TITLE: Weak binding of VX-478 to human plasma proteins and implications for anti-human immunodeficiency virus therapy [see comments]. COMMENT: Comment in: J Infect Dis 1996 Jun; 173(6):1524-6 AUTHOR: Livington D J; Pazhanisamy S; Porter D J; Partaledis J A; Tung R D; Painter G R Vertex Pharmaceuticals Inc., Cambridge, MA 02139, CORPORATE SOURCE: USA. JOURNAL OF INFECTIOUS DISEASES, (1995 Nov) 172 (5) SOURCE: 1238-45. Journal code: IH3. ISSN: 0022-1899. PUB. COUNTRY: United States Journal; Article; (JOURNAL ARTICLE) LANGUAGE: English FILE SEGMENT: Abridged Index Medicus Journals; Priority Journals ENTRY MONTH: 9602 VX-478 is a potent inhibitor of human immunodeficiency virus type 1 (HIV-1) protease (Ki, 0.6 nM) and of HIV-1 replication in antiviral assays (IC90, 80 nM). The fractional binding of VX-478 to human plasma and to purified plasma proteins was determined by equilibrium dialysis and difference UV spectrophotometry. Binding to alpha 1-acid glycoprotein (89% at 2 microM total drug concentration, Kd of 4 microM) accounts for its fractional binding in plasma (93%). Stopped-flow spectrophotometry methods showed that binding is a reversible two-step process. The measured dissociation rate constant

approaches 100 s-1. The antiviral effect of vx-478 was determined in the presence of 45% human plasma, in which the IC90 increased by 1.5-fold compared with control experiments in the presence of 15% fetal bovine serum. The effects of protein binding on the antiviral activity of VX-478 are minor, as expected for a weak drug-protein interaction.

Weak binding of **VX-478** to human plasma proteins TΙ and implications for anti-human immunodeficiency virus therapy [see comments].

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CTCheck Tags: Animal; Human

*Acquired Immunodeficiency Syndrome: DT, drug therapy

*Antiviral Agents: BL, blood

Blood

*Blood Proteins: ME, metabolism

Cattle Fetus

*HIV Protease Inhibitors: BL, blood

Kinetics

Molecular Structure

*Orosomucoid: ME, metabolism

Protein Binding

Spectrophotometry, Ultraviolet

*Sulfonamides: BL, blood

0 (Antiviral Agents); 0 (Blood Proteins); 0 (HIV CN

Protease Inhibitors); 0 (Orosomucoid); 0

(Sulfonamides); 0 (VX 478)

L30 ANSWER 22 OF 30 MEDLINE

ACCESSION NUMBER: 95363927

MEDLINE

TITLE: In vitro selection and characterization of human

immunodeficiency virus type 1 (HIV-1) isolates with reduced sensitivity to hydroxyethylamino sulfonamide

inhibitors of HIV-1 aspartyl protease.

Partaledis J A; Yamaguchi K; Tisdale M; Blair E E; AUTHOR:

Falcione C; Maschera B; Myers R E; Pazhanisamy S;

Futer O; Cullinan A B; et al

Vertex Pharmaceuticals Incorporated, Cambridge, CORPORATE SOURCE:

Massachusetts 02139-4211, USA.

JOURNAL OF VIROLOGY, (1995 Sep) 69 (9) 5228-35. SOURCE:

Journal code: KCV. ISSN: 0022-538X.

PUB. COUNTRY: United States

Journal; Article; (JOURNAL ARTICLE)

LANGUAGE:

AB

English

FILE SEGMENT:

Priority Journals; Cancer Journals

ENTRY MONTH:

9511 Human immunodeficiency virus type 1 (HIV-1) variants with reduced

sensitivity to the hydroxyethylamino sulfonamide protease inhibitors VB-11,328 and VX-478 have been selected in vitro

by two independent serial passage protocols with HIV-1 in CEM-SS and

MT-4 cell lines. Virus populations with greater than

100-fold-increased resistance to both inhibitors compared with the parental virus have been obtained. DNA sequence analyses of the

protease genes from VB-11,328- and VX-478

-resistant variants reveal a sequential accumulation of point mutations, with similar resistance patterns occurring for the two inhibitors. The deduced amino acid substitutions in the resistant protease are Leu-10-->Phe, Met-46-->Ile, Ile-47-->Val, and Ile-50-->Val. This is the first observation in HIV protease resistance studies of an Ile-50-->Val mutation, a mutation that appears to arise uniquely against the sulfonamide inhibitor class. When the substitutions observed were introduced as single mutations into an HIV-1 infectious clone (HXB2), only the Ile-50-->Val mutant showed reduced sensitivity (two- to threefold) to VB-11,328 and

VX-478. A triple protease mutant infectious clone

carrying the mutations Met-46-->Ile, Ile-47-->Val, and Ile-50-->Val, however, showed much greater reduction in sensitivity (14- to

20-fold) to VB-11,328 and **VX-478**. The same

mutations were studied in recombinant HIV protease. The mutant protease Ile-50-->Val displays a much lower affinity for the inhibitors than the parent enzyme (< or = 80-fold). The protease triply mutated at Met-46-->Ile, Ile-47-->Val, and Ile-50-->Val shows an even greater decrease in inhibitor binding (< or = 270-fold). The sulfonamide-resistant HIV protease variants remain sensitive to inhibitors from other chemical classes (Ro 31-8959 and L-735,524),

suggesting possibilities for clinical use of HIV protease inhibitors in combination or serially.

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M46I/V82F, M46I/I84V, and V32I/K45I/F53L/A71V/I84V/L89M--were

generated on the basis of results of in vitro selection experiments using the inhibitors A-77003, A-84538, and KNI-272. Kinetic parameters of mutant and wild-type (WT) enzymes were measured along with inhibition constants (Ki) toward the inhibitors A-77003, A-84538, KNI-272, L-735,524, and Ro31-8959. The catalytic efficiency, kcat/Km, for the mutants decreased relative to WT by a factor of 1.2-14.8 and was mainly due to the elevation of Km. The effects of specific mutations on Ki values were unique with respect to both inhibitor and mutant enzyme. A new property, termed vitality, defined as the ratio (Kikcat/Km) mutant/(Kikcat/Km) WT was introduced to compare the selective advantage of different mutants in the presence of a given inhibitor. High vitality values were generally observed with mutations that emerged during in vitro selection studies. The kinetic model along with the panel of mutants described here should be useful for evaluating and predicting patterns of resistance for HIV PR inhibitors and may aid in the selection of inhibitor combinations to combat drug resistance.

AB Eleven different recombinant, drug-resistant HIV-1 protease (HIV PR) mutants--R8Q, V32I, M46I, V82A, V82F, V82I, I84V, V32I/I84V, M46I/V82F, M46I/I84V, and V32I/K45I/F53L/A71V/I84V/L89M--were generated on the basis of results of in vitro selection experiments using the inhibitors A-77003, A-84538, and KNI-272. Kinetic parameters of mutant and wild-type (WT) enzymes were measured along with inhibition constants (Ki) toward the inhibitors A-77003, A-84538, KNI-272, L-735,524, and Ro31-8959. The catalytic efficiency, kcat/Km, for the mutants decreased relative to WT by a factor of 1.2-14.8 and was mainly due to the elevation of Km. The effects of specific mutations on Ki values were unique with respect to both inhibitor and mutant enzyme. A new property, termed vitality, defined as the ratio (Kikcat/Km) mutant/(Kikcat/Km) WT was introduced to compare the selective advantage of different mutants in the presence of a given inhibitor. High vitality values were generally observed with mutations that emerged during in vitro selection studies. The kinetic model along with the panel of mutants described here should be useful for evaluating and predicting patterns of resistance for HIV PR inhibitors and may aid in the selection of inhibitor combinations to combat drug resistance.

CT Check Tags: Comparative Study

Amino Acid Sequence

Binding Sites

Carbamates: PD, pharmacology

Cloning, Molecular

Drug Resistance, Microbial

*HIV Protease: ME, metabolism

*HIV Protease Inhibitors: PD, pharmacology

*HIV-1: EN, enzymology

Isoquinolines: PD, pharmacology

Kinetics

Methylurea Compounds: PD, pharmacology

Mutagenesis, Site-Directed

Oligopeptides: PD, pharmacology

*Point Mutation

Pyridines: PD, pharmacology Quinolines: PD, pharmacology

Recombinant Proteins: ME, metabolism

Structure-Activity Relationship

11/12/97

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L30 ANSWER 1 OF 30 EMBASE COPYRIGHT 1997 ELSEVIER SCI. B.V.

97299931 EMBASE ACCESSION NUMBER:

TITLE: Human immunodeficiency virus protease inhibitors From

drug design to clinical studies.

AUTHOR: Lin J.H.

J.H. Lin, Drug Metabolism, Mercke Research CORPORATE SOURCE:

Laboratories, West Point, PA 19486, United States

Advanced Drug Delivery Reviews, (1997) 27/2-3 SOURCE:

> (215-233). Refs: 58

ISSN: 0169-409X CODEN: ADDREP

s 0169-409X(97)00044-6 PUBLISHER IDENT .:

Netherlands COUNTRY:

DOCUMENT TYPE: Journal

Pharmacology 030 FILE SEGMENT:

> 037 Drug Literature Index

LANGUAGE: English SUMMARY LANGUAGE: English

The discovery of human immunodeficiency virus (HIV)

protease inhibitors is an example in which

pharmacokinetic evaluation was implemented early in the discovery phase to obtain optimal pharmacological and pharmacokinetic

properties. Currently, three HIV protease

inhibitors, saquinavir, indinavir and ritonavir are

clinically available. As a family, these HIV

protease inhibitors are characterized

pharmacologically by their ability to inhibit the viral protease enzyme. Pharmacokinetically, they are quite different due to their dissimilarity in physicochemical properties. Bioavailability appears to be limited with saquinavir, but not with indinavir and ritonavir. Although all three drugs are metabolized extensively by

cytochrome P-450, saquinavir and indinavir are high clearance drugs while ritonavir is a low clearance drug. Despite their significant differences in elimination clearance, all three HIV proteases are given at high oral doses (600-800 mg) either twice or three times daily. These HIV protease inhibitors

show superior therapeutic activity and a more favorable safety profile than those reported for the established reverse transcriptase inhibitors. However, the potential for interactions with other drugs metabolized by the CYP 3A4 isoform appears to be considerable. In addition, repeated administration of enzyme inducers results in a substantial decrease of plasma concentrations of protease inhibitors. Therefore, co-administration of drugs, such as rifampicin and rifabutin, must be avoided.

HIV protease inhibitors are promising in

the treatment of AIDS. Although they are not a cure, they can significantly inhibit that viral replication and improve the quality of life for people who have HIV infection.

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     EMTAGS: infection (0310); pharmacokinetics (0194); dog (0711);
CT
     mammal (0738); monkey (0725); human (0888); nonhuman (0777); rat
     (0733); oral drug administration (0181); review (0001); priority
     journal (0007)
     Medical Descriptors:
     *human immunodeficiency virus infection
     *pharmacokinetics
     enzyme induction
     drug design
     physical chemistry
     drug bioavailability
     drug absorption
     dog
     monkey
     drug transport
     drug metabolism
     drug protein binding
     human
     nonhuman
     rat
     oral drug administration
     review
     priority journal
     Drug Descriptors:
     *saquinavir: IT, drug interaction
     *saquinavir: PK, pharmacokinetics
     *indinavir: IT, drug interaction
     *indinavir: PK, pharmacokinetics
     *ritonavir: IT, drug interaction
     *ritonavir: PK, pharmacokinetics
     proteinase inhibitor: PK, pharmacokinetics
     zidovudine: IT, drug interaction
     zalcitabine: IT, drug interaction
     rifampicin: IT, drug interaction
     rifabutin: EC, endogenous compound
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clarithromycin: IT, drug interaction stavudine: IT, drug interaction desipramine: IT, drug interaction nifedipine: IT, drug interaction terfenadine: IT, drug interaction dextromethorphan: IT, drug interaction a 77003: PK, pharmacokinetics a 80987: PK, pharmacokinetics (saquinavir) 127779-20-8; (indinavir) 150378-17-9, 157810-81-6; RN (ritonavir) 155213-67-5; (proteinase inhibitor) 37205-61-1; (zidovudine) 30516-87-1; (zalcitabine) 7481-89-2; (rifampicin) 13292-46-1; (rifabutin) 72559-06-9; (clarithromycin) 81103-11-9; (stavudine) 3056-17-5; (desipramine) 50-47-5, 58-28-6; (nifedipine) 21829-25-4; (terfenadine) 50679-08-8; (dextromethorphan) 125-69-9, 125-71-3; (a 77003) 134878-17-4 A 77003; A 80987 CN L30 ANSWER 2 OF 30 EMBASE COPYRIGHT 1997 ELSEVIER SCI. B.V. 97159073 EMBASE ACCESSION NUMBER: TITLE: Hepatic and intestinal metabolism of indinavir, an HIV protease inhibitor, in rat and human microsomes: Major role of CYP3A. Chiba M.; Hensleigh M.; Lin J.H. AUTHOR: Dr. M. Chiba, Merck Research Laboratories, Department CORPORATE SOURCE: of Drug Metabolism, West Point, PA 19486, United Biochemical Pharmacology, (1997) 53/8 (1187-1195). SOURCE: Refs: 22 ISSN: 0006-2952 CODEN: BCPCA6 s 0006-2952(97)00100-7 PUBLISHER IDENT .: United States COUNTRY: DOCUMENT TYPE: Journal FILE SEGMENT: 030 Pharmacology 037 Drug Literature Index LANGUAGE: English SUMMARY LANGUAGE: English The metabolism of indinavir, a human immune deficiency virus (HIV) protease inhibitor, has been characterized extensively in rats and humans. All oxidative metabolites found in vivo were formed when indinavir was incubated with NADPH-fortified hepatic and intestinal microsomes obtained from rats and humans. In vitro kinetic studies revealed that V(max)/K(m) values (.mu.L/min/mg protein) in rat and human liver microsomes were approximately 8- and 2-fold greater than those in the intestinal microsomes of the corresponding species (55.8 and 6.7 for the liver and intestine, respectively, in rats; 16.5 and 7.7 for the liver and intestine, respectively, in humans). However, when V(max)/K(m) was scaled up to intrinsic clearance (mL/min/kg body weight), hepatic intrinsic clearance was much greater than the

intestinal clearance by 50- to 200-fold. These results suggest that

the involvement of CYP3A isoforms in both organs. Oral treatment of rats with dexamethasone (50 mg/kg/day for 4 days), a potent CYP3A

the liver plays a much greater role in first-pass metabolism of indinavir than the intestine in both species. Consistently, ketoconazole, a selective inhibitor for CYP3A, and an anti-rat CYP3A1 antibody strongly inhibited hepatic and intestinal metabolism of indinavir in both rats and humans, suggesting

inducer, increased both hepatic and intestinal **metabolism** of indinavir by a factor of 7 and 3, respectively. Furthermore, indinavir selectively inhibited 6.beta.-hydroxylase activity of testosterone, a GYP3A marker activity, in rat and human liver microsomes; the interactions between testosterone and indinavir were competitive with K(i) values of $< 1.0\,$.mu.M.

TI Hepatic and intestinal metabolism of indinavir, an HIV protease inhibitor, in rat and human microsomes: Major role of CYP3A.

The metabolism of indinavir, a human immune deficiency virus (HIV) protease inhibitor, has been characterized extensively in rats and humans. All oxidative metabolites found in vivo were formed when indinavir was incubated with NADPH-fortified hepatic and intestinal microsomes obtained from rats and humans. In vitro kinetic studies revealed that V(max)/K(m) values (.mu.L/min/mg protein) in rat and human liver microsomes were approximately 8- and 2-fold greater than those in the intestinal microsomes of the corresponding species (55.8 and 6.7 for the liver and intestine, respectively, in rats; 16.5 and 7.7 for the liver and intestine, respectively, in humans). However, when V(max)/K(m) was scaled up to intrinsic clearance (mL/min/kg body weight), hepatic intrinsic clearance was much greater than the intestinal clearance by 50- to 200-fold. These results suggest that the liver plays a much greater role in first-pass metabolism of indinavir than the intestine in both species. Consistently, ketoconazole, a selective inhibitor for CYP3A, and an anti-rat CYP3A1 antibody strongly inhibited hepatic and intestinal metabolism of indinavir in both rats and humans, suggesting the involvement of CYP3A isoforms in both organs. Oral treatment of rats with dexamethasone (50 mg/kg/day for 4 days), a potent CYP3A inducer, increased both hepatic and intestinal metabolism of indinavir by a factor of 7 and 3, respectively. Furthermore, indinavir selectively inhibited 6.beta.-hydroxylase activity of testosterone, a GYP3A marker activity, in rat and human liver microsomes; the interactions between testosterone and indinavir were competitive with K(i) values of < 1.0 .mu.M.

CT EMTAGS: pharmacokinetics (0194); digestive system (0935); liver (0946); mammal (0738); human (0888); nonhuman (0777); male (0041); rat (0733); controlled study (0197); human tissue, cells or cell components (0111); animal tissue, cells or cell components (0105); adolescent (0017); article (0060); priority journal (0007); enzyme (0990)

Medical Descriptors:

AB

*drug metabolism
liver metabolism
intestine
drug oxidation
liver microsome
kinetics
drug clearance
first pass effect
human
nonhuman
male
rat
controlled study
human tissue
animal tissue

adolescent article priority journal Drug Descriptors: *indinavir: PK, pharmacokinetics *proteinase inhibitor: PK, pharmacokinetics reduced nicotinamide adenine dinucleotide phosphate cytochrome p450 isoenzyme testosterone dexamethasone oxygenase CN (1) Mk 639 L30 ANSWER 3 OF 30 EMBASE COPYRIGHT 1997 ELSEVIER SCI. B.V. 97092491 EMBASE ACCESSION NUMBER: Indinavir. TITLE: AUTHOR: Ohta Y.; Shinkai I. CORPORATE SOURCE: Japan Bioorganic and Medicinal Chemistry, (1997) 5/3 SOURCE: (463-464). ISSN: 0968-0896 CODEN: BMECEP PUBLISHER IDENT .: s 0968-0896(96)00261-1 United Kingdom COUNTRY: DOCUMENT TYPE: Journal FILE SEGMENT: 004 Microbiology 030 Pharmacology 037 Drug Literature Index LANGUAGE: English English SUMMARY LANGUAGE: The IC95 (95% inhibitory concentration) of indinavir was in the range of 25-100 nM. In drug combination studies with the nucleoside analogues zidovudine and didanosine, as well as with an investigational non-nucleoside (L-697,661), indinavir showed synergistic activity in cell culture. Viral resistance was correlated with the accumulation of mutations that resulted in the expression of amino acid substitutions in the viral protease. Eleven amino acid residue positions have been identified. Indinavir was rapidly absorbed in the fasted state. Cross-resistance was noted between indinavir and the protease inhibitor ritonavir. Varying degrees of cross-resistance have been observed between indinavir and other HIV-protease inhibitors. Seven metabolites have been identified, one glucuronide conjugate and six oxidation metabolites. In vitro studies indicate that cytochrome P-450 3A4 (CRY3A4) is the major enzyme responsible for formation of the oxidative metabolites. Indinavir has been studied in phase III clinical trials as a monotherapy (dose-escalation) and in combination with zidovudine and with zidovudine + didanosine. The recommended dosage of Crixivan is 800 mg (two 400 mg capsules) orally every 8 h. The dosage is the same whether Crixivan is used alone or in combination with other antiretroviral agents. In an analysis of early clinical trials for safety, nephrolithiasis was the only clinically significant ADR. The IC95 (95% inhibitory concentration) of indinavir was in the range of 25-100 nM. In drug combination studies with the nucleoside analogues zidovudine and didanosine, as well as with an investigational non-nucleoside (L-697,661), indinavir showed synergistic activity in cell culture. Viral resistance was

correlated with the accumulation of mutations that resulted in the

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degrees of cross-resistance have been observed between indinavir and
other HIV-protease inhibitors. Seven
metabolites have been identified, one glucuronide conjugate
and six oxidation metabolites. In vitro studies indicate
that cytochrome P-450 3A4 (CRY3A4) is the major enzyme responsible
for formation of the oxidative metabolites. Indinavir has
been studied in phase III clinical trials as a monotherapy
(dose-escalation) and in combination with zidovudine and with
zidovudine + didanosine. The recommended dosage of Crixivan is 800
mg (two 400 mg capsules) orally every 8 h. The dosage is the same
whether Crixivan is used alone or in combination with other
antiretroviral agents. In an analysis of early clinical trials for
safety, nephrolithiasis was the only clinically significant ADR.
EMTAGS: infection (0310); therapy (0160); virus (0761); cell, tissue
or organ culture (0103); chemical procedures (0107); heredity
(0137); pharmacokinetics (0194); mammal (0738); human (0888); oral
drug administration (0181); short survey (0002)
Medical Descriptors:
*antiviral activity
*human immunodeficiency virus infection: DT, drug therapy
*human immunodeficiency virus
in vitro study
cell culture
drug synthesis
drug resistance
mutation
drug metabolism
human
oral drug administration
short survey
Drug Descriptors:
*indinavir: AN, drug analysis
*indinavir: CB, drug combination
*indinavir: CM, drug comparison
*indinavir: DV, drug development
*indinavir: DO, drug dose
*indinavir: DT, drug therapy
*indinavir: PD, pharmacology
*antiretrovirus agent: AN, drug analysis
*antiretrovirus agent: CB, drug combination
*antiretrovirus agent: CM, drug comparison
*antiretrovirus agent: DV, drug development
*antiretrovirus agent: IT, drug interaction
*antiretrovirus agent: PD, pharmacology
zidovudine: CB, drug combination
zidovudine: DT, drug therapy
didanosine: CB, drug combination
didanosine: DT, drug therapy
3 [(4,7 dichloro 2 benzoxazolylmethyl)amino] 5 ethyl 6 methyl 2(1h)
pyridone: CB, drug combination
3 [(4,7 dichloro 2 benzoxazolylmethyl)amino] 5 ethyl 6 methyl 2(1h)
pyridone: DT, drug therapy
drug metabolite: AN, drug analysis
(1) Crixivan; (2) Mk 639; (3) L 735524; (4) L 697661
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CN

CT

L30 ANSWER 4 OF 30 EMBASE COPYRIGHT 1997 ELSEVIER SCI. B.V. 97069869 EMBASE ACCESSION NUMBER: Pharmacokinetic enhancement of inhibitors of the TITLE: human immunodeficiency virus protease by coadministration with ritonavir. Kempf D.J.; Marsh K.C.; Kumar G.; Rodrigues A.D.; AUTHOR: Denissen J.F.; McDonald E.; Kukulka M.J.; Hsu A.; Granneman G.R.; Baroldi P.A.; Sun E.; Pizzuti D.; Plattner J.J.; Norbeck D.W.; Leonard J.M. United States. Dale.J.Kempf@abbott.com CORPORATE SOURCE: SOURCE: Antimicrobial Agents and Chemotherapy, (1997) 41/3 (654-660).Refs: 33 ISSN: 0066-4804 CODEN: AMACCQ United States COUNTRY: DOCUMENT TYPE: Journal FILE SEGMENT: 004 Microbiology Pharmacology 030 037 Drug Literature Index English LANGUAGE: SUMMARY LANGUAGE: English Coadministration with the human immunodeficiency virus (HIV) protease inhibitor ritonavir was investigated as a method for enhancing the levels of other peptidomimetic HIV protease inhibitors in plasma. In rat and human liver microsomes, ritonavir potently inhibited the cytochrome P450 (CYP) - mediated metabolism of saquinavir, indinavir, nelfinavir, and VX-478. The structural features of ritonavir responsible for CYP binding and inhibition were examined. Coadministration of other protease inhibitors with ritonavir in rats and dogs produced elevated and sustained plasma drug levels 8 to 12 h after a single dose. Drug exposure in rats was elevated by 8- to 46-fold. A >50-fold enhancement of the concentrations of saquinavir in plasma was observed in humans following a single codose of ritonavir (600 mg) and saquinavir (200 mg). These results indicate that ritonavir can favorably alter the pharmacokinetic profiles of other protease inhibitors. Combination regimens of ritonavir and other protease inhibitors may thus play a role in the treatment of HIV infection. Because of potentially substantial drug level increases, however, such combinations require further investigation to establish safe regimens for clinical use. Coadministration with the human immunodeficiency virus (HIV AB) protease inhibitor ritonavir was investigated as a method for enhancing the levels of other peptidomimetic HIV protease inhibitors in plasma. In rat and human liver microsomes, ritonavir potently inhibited the cytochrome P450 (CYP) - mediated metabolism of saquinavir, indinavir, nelfinavir, and VX-478. The structural features of ritonavir responsible for CYP binding and inhibition were examined. Coadministration of other protease inhibitors with ritonavir in rats and dogs produced elevated and sustained plasma drug levels 8 to 12 h after a single dose. Drug exposure in rats was elevated by 8- to 46-fold. A >50-fold

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     inhibitors. Combination regimens of ritonavir and other protease
     inhibitors may thus play a role in the treatment of HIV infection.
     Because of potentially substantial drug level increases, however,
     such combinations require further investigation to establish safe
     regimens for clinical use.
    EMTAGS: virus (0761); infection (0310); therapy (0160); etiology
CT
     (0135); digestive system (0935); liver (0946); pharmacokinetics
     (0194); mammal (0738); human (0888); nonhuman (0777); male (0041);
     female (0042); rat (0733); human experiment (0104); normal human
     (0800); animal experiment (0112); human tissue, cells or cell
     components (0111); oral drug administration (0181); article (0060);
     priority journal (0007)
    Medical Descriptors:
     *human immunodeficiency virus 1
     *human immunodeficiency virus infection: DT, drug therapy
     *human immunodeficiency virus infection: ET, etiology
     liver microsome
     drug metabolism
     dose response
     drug mixture
     drug elimination
     drug potentiation
     human
     nonhuman
     male
     female
     rat
     human experiment
     normal human
     clinical trial
     crossover procedure
     animal experiment
     human tissue
     oral drug administration
     article
     priority journal
     Drug Descriptors:
     *ritonavir: CT, clinical trial
     *ritonavir: CB, drug combination
     *ritonavir: DO, drug dose
     *ritonavir: IT, drug interaction
     *ritonavir: DT, drug therapy
     *ritonavir: PK, pharmacokinetics
     *ritonavir: PD, pharmacology
     *proteinase inhibitor: CT, clinical trial
     *proteinase inhibitor: CB, drug combination
     *proteinase inhibitor: DO, drug dose
     *proteinase inhibitor: IT, drug interaction
     *proteinase inhibitor: DT, drug therapy
     *proteinase inhibitor: PK, pharmacokinetics
     *proteinase inhibitor: PD, pharmacology
     saguinavir: CT, clinical trial
     saguinavir: CB, drug combination
     saquinavir: DO, drug dose
     saquinavir: IT, drug interaction
     saquinavir: DT, drug therapy
     saquinavir: PK, pharmacokinetics
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saquinavir: PD, pharmacology
     indinavir: CB, drug combination
     indinavir: DV, drug development
     indinavir: DO, drug dose
     indinavir: IT, drug interaction
     indinavir: DT, drug therapy
     indinavir: PK, pharmacokinetics
     indinavir: PD, pharmacology
     nelfinavir: CB, drug combination
     nelfinavir: DV, drug development
nelfinavir: DO, drug dose
     nelfinavir: IT, drug interaction
nelfinavir: DT, drug therapy
     nelfinavir: PK, pharmacokinetics
     nelfinavir: PD, pharmacology
     4 amino n [2 hydroxy 4 phenyl 3 (tetrahydrofuran 3
     yloxycarbonylamino)butyl] n isobutylbenzenesulfonamide: CB, drug
     combination
     4 amino n [2 hydroxy 4 phenyl 3 (tetrahydrofuran 3
     yloxycarbonylamino)butyl] n isobutylbenzenesulfonamide: DV, drug
     development
     4 amino n [2 hydroxy 4 phenyl 3 (tetrahydrofuran 3
     yloxycarbonylamino)butyl] n isobutylbenzenesulfonamide: DO, drug
     dose
     4 amino n [2 hydroxy 4 phenyl 3 (tetrahydrofuran 3
     yloxycarbonylamino)butyl] n isobutylbenzenesulfonamide: IT, drug
     interaction
     4 amino n [2 hydroxy 4 phenyl 3 (tetrahydrofuran 3
     yloxycarbonylamino)butyl] n isobutylbenzenesulfonamide: DT, drug
     therapy
     4 amino n [2 hydroxy 4 phenyl 3 (tetrahydrofuran 3
     yloxycarbonylamino)butyl] n isobutylbenzenesulfonamide: PK,
     pharmacokinetics
     4 amino n [2 hydroxy 4 phenyl 3 (tetrahydrofuran 3
     yloxycarbonylamino)butyl] n isobutylbenzenesulfonamide: PD,
     pharmacology
     cytochrome p450
     Vx 478
L30 ANSWER 5 OF 30 EMBASE COPYRIGHT 1997 ELSEVIER SCI. B.V.
                     97060069 EMBASE
ACCESSION NUMBER:
                     Selective biotransformation of the human
TITLE:
                     immunodeficiency virus protease inhibitor saquinavir
                    by human small-intestinal cytochrome P4503A4:
                     Potential contribution to high first-pass
                  metabolism.
                     Fitzsimmons M.E.; Collins J.M.
AUTHOR:
CORPORATE SOURCE:
                     United States
                     Drug Metabolism and Disposition, (1997) 25/2
SOURCE:
                     (256-266).
                     Refs: 44
                     ISSN: 0090-9556 CODEN: DMDSAI
                     United States
COUNTRY:
DOCUMENT TYPE:
                     Journal
FILE SEGMENT:
                     030
                             Pharmacology
                             Drug Literature Index
                     037
                     English
LANGUAGE:
                    English
SUMMARY LANGUAGE:
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CN

Saquinavir is a HIV1 protease inhibitor used in the treatment of AB patients with acquired immunodeficiency syndrome, but its use is limited by low oral bioavailability. The potential of human intestinal tissue to metabolize saquinavir was assessed in 17 different human smell-intestinal microsomal preparations. Saquinavir was metabolized by human small-intestinal microsomes to numerous mono- and dihydroxylated species with K(M) values of 0.3-0.5 .mu.M. The major metabolites M-2 and M-7 were single hydroxylations on the octahydro-2-(1H)-isoquinolinyl and (1,1-dimethylethyl)amino groups, respectively. Ketoconazole and troleandomycin, selective inhibitors of cytochrome P4503A4 (CYP3A4), were potent inhibitors for all oxidative metabolites of saquinavir. The cytochrome P450-selective inhibitors furafylline, fluvoxamine, sulfaphenazole, mephenytoin, quinidine, and chlorzoxazone had little inhibitory effect. All saquinavir metabolites were highly correlated with testosterone 6.beta.-hydroxylation and with each other. Human hepatic microsomes and recombinant CYP3A4 oxidized saquinavir to the same metabolic profile observed with human small-intestinal microsomes. Indinavir, a potent HIV protease inhibitor and a substrate for human hepatic CYP3A4, was a comparatively poor substrate for human intestinal microsomes and inhibited the oxidative metabolism of saquinavir to all metabolites with a K(i) of 0.2 .mu.M. In addition, saquinavir inhibited the human, small- intestinal, microsomal CYP3A4-dependent detoxication pathway of terfenadine to its alcohol metabolite with a K(i) value of 0.7 .mu.M. These data indicate that saquinavir is metabolized by human intestinal CYP3A4, that this metabolism may contribute to its poor oral bioavailability, and that combination therapy with indinavir or other protease inhibitors may attenuate its low relative bioavailability.

- TI Selective biotransformation of the human immunodeficiency virus protease inhibitor saquinavir by human small-intestinal cytochrome P4503A4: Potential contribution to high first-pass metabolism.
- Saquinavir is a HIV1 protease inhibitor used in the treatment of AB patients with acquired immunodeficiency syndrome, but its use is limited by low oral bioavailability. The potential of human intestinal tissue to metabolize saquinavir was assessed in 17 different human smell-intestinal microsomal preparations. Saquinavir was metabolized by human small-intestinal microsomes to numerous mono- and dihydroxylated species with K(M) values of 0.3-0.5 .mu.M. The major metabolites M-2 and M-7 were single hydroxylations on the octahydro-2-(1H)-isoquinolinyl and (1,1-dimethylethyl)amino groups, respectively. Ketoconazole and troleandomycin, selective inhibitors of cytochrome P4503A4 (CYP3A4), were potent inhibitors for all oxidative metabolites of saquinavir. The cytochrome P450-selective inhibitors furafylline, fluvoxamine, sulfaphenazole, mephenytoin, quinidine, and chlorzoxazone had little inhibitory effect. All saquinavir metabolites were highly correlated with testosterone 6.beta.-hydroxylation and with each other. Human hepatic microsomes and recombinant CYP3A4 oxidized saquinavir to the same metabolic profile observed with human small-intestinal microsomes. Indinavir, a potent HIV protease inhibitor and a substrate for human hepatic CYP3A4, was a comparatively poor substrate for human intestinal microsomes and

```
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     CYP3A4-dependent detoxication pathway of terfenadine to its alcohol
     metabolite with a K(i) value of 0.7 .mu.M. These data
     indicate that saquinavir is metabolized by human
     intestinal CYP3A4, that this metabolism may contribute to
     its poor oral bioavailability, and that combination therapy with
     indinavir or other protease inhibitors may attenuate its low
     relative bioavailability.
     EMTAGS: pharmacokinetics (0194); digestive system (0935); small
CT
     intestine (0941); liver (0946); mammal (0738); human (0888);
     controlled study (0197); human tissue, cells or cell components
     (0111); article (0060); priority journal (0007); enzyme (0990)
     Medical Descriptors:
     *biotransformation
     *drug metabolism
     small intestine
     microsome
     liver microsome
     enzyme inhibition
     human
     controlled study
     human tissue
     article
     priority journal
     Drug Descriptors:
     *proteinase inhibitor: PK, pharmacokinetics
     *saquinavir: PK, pharmacokinetics
     *indinavir: PK, pharmacokinetics
     *cytochrome p450 isoenzyme: EC, endogenous compound
     *cytochrome p450 inhibitor
     alpha naphthoflavone
     furafylline
     fluvoxamine
     quercetin
     sulfaphenazole
     mephenytoin
     quinidine
     chlorzoxazone
     ketoconazole
     troleandomycin
     midazolam
     cyclosporin a
     terfenadine
     (1) Ro 31 8959; (2) Mk 639
L30 ANSWER 6 OF 30 EMBASE COPYRIGHT 1997 ELSEVIER SCI. B.V.
ACCESSION NUMBER:
                     97033183 EMBASE
                     Indinavir: A pharmacologic and clinical review of a
TITLE:
                     new HIV protease
                   inhibitor.
                     Lacy M.K.; Abriola K.P.
AUTHOR:
CORPORATE SOURCE:
                     United States
                     Connecticut Medicine, (1996) 60/12 (723-727).
SOURCE:
                     Refs: 20
                     ISSN: 0010-6178 CODEN: CNMEAH
```

United States

COUNTRY:

Journal

DOCUMENT TYPE:

FILE SEGMENT: 004 Microbiology 030 Pharmacology Drug Literature Index 037 LANGUAGE: English Indinavir: A pharmacologic and clinical review of a new HIV protease inhibitor. EMTAGS: infection (0310); pharmacokinetics (0194); therapy (0160); CTmammal (0738); human (0888); short survey (0002) Medical Descriptors: *human immunodeficiency virus infection drug information drug mechanism drug metabolism drug indication human short survey Drug Descriptors: *indinavir: IT, drug interaction *indinavir: PD, pharmacology *indinavir: PK, pharmacokinetics *proteinase inhibitor: PD, pharmacology
*proteinase inhibitor: PK, pharmacokinetics
*proteinase inhibitor: IT, drug interaction zidovudine: DT, drug therapy zidovudine: IT, drug interaction lamivudine: IT, drug interaction stavudine: IT, drug interaction stavudine: DT, drug therapy cimetidine: IT, drug interaction quinidine: IT, drug interaction cotrimoxazole: IT, drug interaction fluconazole: IT, drug interaction isoniazid: IT, drug interaction clarithromycin: IT, drug interaction rifampicin: IT, drug interaction didanosine: IT, drug interaction terfenadine: IT, drug interaction astemizole: IT, drug interaction cisapride: IT, drug interaction triazolam: IT, drug interaction midazolam: IT, drug interaction ketoconazole: IT, drug interaction (1) Crixivan; (2) Mk 639; (3) L 735524 L30 ANSWER 7 OF 30 MEDLINE 97209067 MEDLINE ACCESSION NUMBER: Pharmacokinetic enhancement of inhibitors of the TITLE: human immunodeficiency virus protease by coadministration with ritonavir. Kempf D J; Marsh K C; Kumar G; Rodrigues A D; AUTHOR: Denissen J F; McDonald E; Kukulka M J; Hsu A; Granneman G R; Baroldi P A; Sun E; Pizzuti D; Plattner J J; Norbeck D W; Leonard J M Department of Infectious Diseases Research, Abbott CORPORATE SOURCE: Laboratories, Illinois 60064, USA... Dale.J.Kempf@abbott.com ANTIMICROBIAL AGENTS AND CHEMOTHERAPY, (1997 Mar) 41 SOURCE:

(3) 654-60.

Journal code: 6HK. ISSN: 0066-4804.

PUB. COUNTRY:

United States
Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

FILE SEGMENT: Priority Journals

ENTRY MONTH: 9708 ENTRY WEEK: 19970801

Coadministration with the human immunodeficiency virus (HIV AB) protease inhibitor ritonavir was investigated as a method for enhancing the levels of other peptidomimetic HIV protease inhibitors in plasma. In rat and human liver microsomes, ritonavir potently inhibited the cytochrome P450 (CYP)-mediated metabolism of saquinavir, indinavir, nelfinavir, and VX-478. The structural features of ritonavir responsible for CYP binding and inhibition were examined. Coadministration of other protease inhibitors with ritonavir in rats and dogs produced elevated and sustained plasma drug levels 8 to 12 h after a single dose. Drug exposure in rats was elevated by 8- to 46-fold. A > 50-fold enhancement of the concentrations of saquinavir in plasma was observed in humans following a single codose of ritonavir (600 mg) and saquinavir (200 mg). These results indicate that ritonavir can favorably alter the pharmacokinetic profiles of other protease inhibitors. Combination regimens of ritonavir and other protease inhibitors may thus play a role in the treatment of HIV infection. Because of potentially substantial drug level increases, however, such combinations require further investigation to establish safe regimens for clinical use.

AB Coadministration with the human immunodeficiency virus (HIV) protease inhibitor ritonavir was investigated as a method for enhancing the levels of other peptidomimetic HIV protease inhibitors in plasma. In rat and human liver microsomes, ritonavir potently inhibited the cytochrome P450 (CYP)-mediated metabolism of saquinavir, indinavir, nelfinavir, and VX-478. The structural features of ritonavir responsible for CYP binding and inhibition were examined. Coadministration of other protease inhibitors with ritonavir in rats and dogs produced elevated and sustained plasma drug levels 8 to 12 h after a single dose. Drug exposure in rats was elevated by 8- to 46-fold. A > 50-fold enhancement of the concentrations of saquinavir in plasma was observed in humans following a single codose of ritonavir (600 mg) and saquinavir (200 mg). These results indicate that ritonavir can favorably alter the pharmacokinetic profiles of other protease inhibitors. Combination regimens of ritonavir and other protease inhibitors may thus play a role in the treatment of HIV infection. Because of potentially substantial drug level increases, however, such combinations require further investigation to establish safe regimens for clinical use.

CT Check Tags: Animal; Female; Human; Male

*Anti-HIV Agents: PD, pharmacology

Area Under Curve

Cytochrome P-450: AI, antagonists & inhibitors

Cytochrome P-450: ME, metabolism

Dogs

Drug Interactions

HIV Protease Inhibitors: PD, pharmacology

*HIV Protease Inhibitors: PK, pharmacokinetics

Rats

Rats, Sprague-Dawley

*Ritonavir: PD, pharmacology

CN 0 (Anti-HIV Agents); 0 (HIV Protease

Inhibitors); 0 (Ritonavir)

L30 ANSWER 8 OF 30 MEDLINE

ACCESSION NUMBER: 97126223 MEDLINE

TITLE: Disposition of indinavir, a potent HIV-1 protease

inhibitor, after an oral dose in humans.

AUTHOR: Balani S K; Woolf E J; Hoagland V L; Sturgill M G;

Deutsch P J; Yeh K C; Lin J H

CORPORATE SOURCE: Department of Drug Metabolism, Merck Research

Laboratories, West Point, PA 19486, USA.

SOURCE: DRUG METABOLISM AND DISPOSITION, (1996 Dec) 24 (12)

1389-94.

Journal code: EBR. ISSN: 0090-9556.

PUB. COUNTRY: United States

(CLINICAL TRIAL)

Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

FILE SEGMENT: Priority Journals

ENTRY MONTH: 9706

ENTRY WEEK: 19970601

Indinavir, N-[2(R)-hydroxy-1(S)-indanyl]-5-[2(S)-tertiarybutylaminocarbonyl-4-(3-pyridylmethyl)piperazino]-4(S)hydroxy-2(R)-phenylmethylpentanamide (L-735,524,MK-639, ayl-4- Crixivan), is a potent and specific inhibitor of the HIV-1(3 protease for the treatment of AIDS. Disposition of [14C]indinavir was investigated in six healthy subjects after single oral administration of 400 mg. AUC, Cmax, and Tmax values for indinavir were 492 microM x min, 4.7 microM, and 50 min, respectively. The AUC value for the total radioactivity in plasma was 1.9 times higher than that of indinavir, indicating the presence of metabolites. The major excretory route was through feces, and the minor through urine. Mean recovery of radioactivity in the feces was 83.4%. In the urine, mean recoveries of the total radioactivity and unchanged indinavir were 18.7% and 11.0% of the dose, respectively. HPLC radioactivity and LC-MS/MS analyses of urine showed the presence of indinavir and low levels of quaternary pyridine N-glucuronide (M1), 2',3'-trans-dihydroxyindanylpyridine N-oxide (M2), 2',3'-trans-dihydroxyindan (M3) and pyridine N-oxide (M4a) analogs, and despyridylmethyl analogs of M3 (M5) and indinavir (M6). M5 and M6 were the major metabolites in urine. The metabolic profile in plasma was similar to that in urine. Quantitatively, the metabolites in feces accounted for >47% of the dose, which along with the urinary excretion of approximately 19%, suggested that the absorption of the drug was appreciable. In the feces, radioactivity was predominantly due to M3, M5, M6, and the parent compound. Thus, in urine and feces, the prominent metabolic pathways were oxidations and oxidative N-dealkylations. Excretion of the quaternary N-glucuronide

AB Indinavir, N-[2(R)-hydroxy-1(S)-indanyl]-5-[2(S)-tertiary-butylaminocarbonyl-4-(3-pyridylmethyl)piperazino]-4(S)-hydroxy-2(R)-phenylmethylpentanamide (L-735,524,MK-

metabolite in the urine, which is a minor metabolite

in human, was specific to primates.

639, ayl-4- Crixivan), is a potent and specific inhibitor of the HIV-1(3 protease for the treatment of AIDS. Disposition of [14C]indinavir was investigated in six healthy subjects after single oral administration of 400 mg. AUC, Cmax, and Tmax values for indinavir were 492 microM x min, 4.7 microM, and 50 min, respectively. The AUC value for the total radioactivity in plasma was 1.9 times higher than that of indinavir, indicating the presence of metabolites. The major excretory route was through feces, and the minor through urine. Mean recovery of radioactivity in the feces was 83.4%. In the urine, mean recoveries of the total radioactivity and unchanged indinavir were 18.7% and 11.0% of the dose, respectively. HPLC radioactivity and LC-MS/MS analyses of urine showed the presence of indinavir and low levels of quaternary pyridine N-glucuronide (M1), 2',3'-trans-dihydroxyindanylpyridine N-oxide (M2), 2',3'-trans-dihydroxyindan (M3) and pyridine N-oxide (M4a) analogs, and despyridylmethyl analogs of M3 (M5) and indinavir (M6). M5 and M6 were the major metabolites in urine. The metabolic profile in plasma was similar to that in urine. Quantitatively, the metabolites in feces accounted for >47% of the dose, which along with the urinary excretion of approximately 19%, suggested that the absorption of the drug was appreciable. In the feces, radioactivity was predominantly due to M3, M5, M6, and the parent compound. Thus, in urine and feces, the prominent metabolic pathways were oxidations and oxidative N-dealkylations. Excretion of the quaternary N-glucuronide metabolite in the urine, which is a minor metabolite in human, was specific to primates.

CT Check Tags: Animal; Female; Human; Male

Adult

Area Under Curve
Bile: ME, metabolism
Biotransformation

Chromatography, High Pressure Liquid

Chromatography, Liquid

Dogs

Feces: CH, chemistry

*HIV Protease Inhibitors: PK, pharmacokinetics

HIV Protease Inhibitors: UR, urine

*HIV-1: EN, enzymology

*Indinavir: PK, pharmacokinetics

Indinavir: UR, urine

Rats

Rats, Sprague-Dawley Species Specificity

Spectrophotometry, Ultraviolet

Spectrum Analysis, Mass

CN 0 (HIV Protease Inhibitors)

L30 ANSWER 9 OF 30 MEDLINE

ACCESSION NUMBER: 97126022 MEDLINE

TITLE:

Genetic correlates of in vivo viral resistance to indinavir, a human immunodeficiency virus type 1

protease inhibitor.

AUTHOR:

Condra J H; Holder D J; Schleif W A; Blahy O M; Danovich R M; Gabryelski L J; Graham D J; Laird D;

Quintero J C; Rhodes A; Robbins H L; Roth E;

Shivaprakash M; Yang T; Chodakewitz J A; Deutsch P J; Leavitt R Y; Massari F E; Mellors J W; Squires K E;

Steigbigel R T; Teppler H; Emini E A

CORPORATE SOURCE: Department of Antiviral Research, Merck Research

Laboratories, West Point, Pennsylvania 19486, USA..

jon condra@merck.com

SOURCE: JOURNAL OF VIROLOGY, (1996 Dec) 70 (12) 8270-6.

Journal code: KCV. ISSN: 0022-538X.

PUB. COUNTRY: United States

Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

FILE SEGMENT: Priority Journals; Cancer Journals OTHER SOURCE: GENBANK-U71606; GENBANK-U72026

ENTRY MONTH: 9703 ENTRY WEEK: 19970304

AB Indinavir (IDV) (also called CRIXIVAN, MK-639,

or L-735,524) is a potent and selective inhibitor of the human immunodeficiency virus type 1 (HIV-1) protease. During early clinical trials, in which patients initiated therapy with suboptimal dosages of IDV, we monitored the emergence of viral resistance to the inhibitor by genotypic and phenotypic characterization of primary HIV-1 isolates. Development of resistance coincided with variable patterns of multiple substitutions among at least 11 protease amino acid residues. No single substitution was present in all resistant isolates, indicating that resistance evolves through multiple genetic pathways. Despite this complexity, all of 29 resistant isolates tested exhibited alteration of residues M-46 (to I or L) and/or V-82 (to A, F, or T), suggesting that screening of these residues may be useful in predicting the emergence of resistance. We also extended our previous finding that IDV-resistant viral variants exhibit various patterns of cross-resistance to a diverse panel of HIV-1 protease inhibitors. Finally, we noted an association between the number of protease amino acid substitutions and the observed level of IDV resistance. No single substitution or pair of substitutions tested gave rise to measurable viral resistance to IDV. The evolution of this resistance was found to be cumulative, indicating the need for ongoing viral replication in this process. These observations strongly suggest that therapy should be initiated with the most efficacious regimen available, both to suppress viral spread and to inhibit the replication that is required for the evolution of resistance.

AΒ Indinavir (IDV) (also called CRIXIVAN, MK-639, or L-735,524) is a potent and selective inhibitor of the human immunodeficiency virus type 1 (HIV-1) protease. During early clinical trials, in which patients initiated therapy with suboptimal dosages of IDV, we monitored the emergence of viral resistance to the inhibitor by genotypic and phenotypic characterization of primary HIV-1 isolates. Development of resistance coincided with variable patterns of multiple substitutions among at least 11 protease amino acid residues. No single substitution was present in all resistant isolates, indicating that resistance evolves through multiple genetic pathways. Despite this complexity, all of 29 resistant isolates tested exhibited alteration of residues M-46 (to I or L) and/or V-82 (to A, F, or T), suggesting that screening of these residues may be useful in predicting the emergence of resistance. We also extended our previous finding that IDV-resistant viral variants exhibit various patterns of cross-resistance to a diverse panel of HIV-1 protease inhibitors. Finally, we noted an association between the number of protease amino acid substitutions and the observed level of IDV resistance. No single substitution or

pair of substitutions tested gave rise to measurable viral resistance to IDV. The evolution of this resistance was found to be cumulative, indicating the need for ongoing viral replication in this process. These observations strongly suggest that therapy should be initiated with the most efficacious regimen available, both to suppress viral spread and to inhibit the replication that is required for the evolution of resistance.

CT Check Tags: Human

Base Sequence

Drug Resistance, Microbial

DNA, Viral Genotype Hela Cells

HIV Infections: DT, drug therapy *HIV Infections: VI, virology HIV Protease: CH, chemistry

*HIV Protease: ME, metabolism

*HIV Protease Inhibitors: PD, pharmacology

HIV-1: CL, classification *HIV-1: DE, drug effects HIV-1: EN, enzymology

HIV-1: IP, isolation & purification

*Indinavir: PD, pharmacology
Molecular Sequence Data

Phenotype

Variation (Genetics)

CN EC 3.4.23.- (HIV Protease); 0 (DNA, Viral); 0 (HIV

Protease Inhibitors)

L30 ANSWER 10 OF 30 MEDLINE

ACCESSION NUMBER:

97125958 MEDLINE

TITLE:

Human immunodeficiency virus. Mutations in the viral

protease that confer resistance to saquinavir increase the dissociation rate constant of the

protease-saquinavir complex.

AUTHOR:

Maschera B; Darby G; Palu G; Wright L L; Tisdale M;

Myers R; Blair E D; Furfine E S

CORPORATE SOURCE:

Department of Virology, Glaxo Wellcome, Stevenage SG1

2NY, United Kingdom.

SOURCE:

JOURNAL OF BIOLOGICAL CHEMISTRY, (1996 Dec 27) 271

(52) 33231-5.

Journal code: HIV. ISSN: 0021-9258.

PUB. COUNTRY:

United States

Journal; Article; (JOURNAL ARTICLE)

LANGUAGE:

English

FILE SEGMENT:

Priority Journals; Cancer Journals

ENTRY MONTH: ENTRY WEEK: 9704 19970401

AB Mutations in the human immunodeficiency virus (HIV) protease (L90M, G48V, and L90M/G48V) arise when HIV is passaged in the presence of the HIV protease inhibitor saquinavir.

These mutations yield a virus with less sensitivity to the drug (L90M > G48V >> L90M/G48V). L90M, G48V, and L90M/G48V proteases have 1/20, 1/160, and 1/1000 the affinity for saquinavir compared to WT protease, respectively. Therefore, the affinity of mutant protease for saquinavir decreased as the sensitivity of the virus to saquinavir decreased. Association rate constants for WT and mutant proteases with saquinavir were similar, ranging from 2 to $4 \times 10(7)$

M-1 s-1. In contrast, the dissociation rate constants for WT, L90M, G48V, and L90M/G48V proteases complexed with saquinavir were 0.0014, 0.019, 0.128, and 0.54 s-1, respectively. This indicated that the reduced affinity for mutant proteases and saquinavir is primarily the result of larger dissociation rate constants. The increased dissociation rate constants may be the result of a decrease in the internal equilibrium between the bound inhibitor with the protease flaps up and the bound inhibitor with the flaps down. Interestingly, the affinity of these mutant proteases for VX-478 , ABT-538, AG-1343, or L-735,524 was not reduced as much as that for saguinavir. Finally, the catalytic constants of WT and mutant proteases were determined for eight small peptide substrates that mimic the viral cleavage sites in vivo. WT and L90M proteases had similar catalytic constants for these substrates. In contrast, G48V and L90M/G48V proteases had catalytic efficiency (kcat/Km) values with TLNF-PISP, RKIL-FLDG, and AETF-YVDG that were 1/10 to 1/20 the value of WT protease. The decreased catalytic efficiencies were primarily the result of increased Km values. Thus, mutations in the protease decrease the affinity of the enzyme for saquinavir and the catalytic efficiency with peptide substrates.

Mutations in the human immunodeficiency virus (HIV) protease (L90M, G48V, and L90M/G48V) arise when HIV is passaged in the presence of the HIV protease inhibitor saquinavir.

These mutations yield a virus with less sensitivity to the drug (L90M > G48V >> L90M/G48V). L90M, G48V, and L90M/G48V proteases have 1/20, 1/160, and 1/1000 the affinity for saquinavir compared to WT protease, respectively. Therefore, the affinity of mutant protease for saquinavir decreased as the sensitivity of the virus to saquinavir decreased. Association rate constants for WT and mutant proteases with saquinavir were similar, ranging from 2 to 4 x 10(7) M-1 s-1. In contrast, the dissociation rate constants for WT, L90M, G48V, and L90M/G48V proteases complexed with saquinavir were 0.0014, 0.019, 0.128, and 0. 54 s-1, respectively. This indicated that the reduced affinity for mutant proteases and saquinavir is primarily the result of larger dissociation rate constants. The increased dissociation rate constants may be the result of a decrease in the internal equilibrium between the bound inhibitor with the protease flaps up and the bound inhibitor with the flaps down. Interestingly, the affinity of these mutant proteases for VX-478

, ABT-538, AG-1343, or L-735,524 was not reduced as much as that for saquinavir. Finally, the catalytic constants of WT and mutant proteases were determined for eight small peptide substrates that mimic the viral cleavage sites in vivo. WT and L90M proteases had similar catalytic constants for these substrates. In contrast, G48V and L90M/G48V proteases had catalytic efficiency (kcat/Km) values with TLNF-PISP, RKIL-FLDG, and AETF-YVDG that were 1/10 to 1/20 the value of WT protease. The decreased catalytic efficiencies were primarily the result of increased Km values. Thus, mutations in the protease decrease the affinity of the enzyme for saquinavir and the catalytic efficiency with peptide substrates.

CT Check Tags: Human

AB

Antiviral Agents: ME, metabolism Drug Resistance, Microbial *HIV Protease: GE, genetics HIV Protease: ME, metabolism

HIV Protease Inhibitors: ME, metabolism

Indinavir: ME, metabolism
Isoquinolines: ME, metabolism

Kinetics Mutagenesis

Ritonavir: ME, metabolism Saquinavir: ME, metabolism *Saquinavir: TU, therapeutic use Sulfonamides: ME, metabolism Sulfonic Acids: ME, metabolism

CN EC 3.4.23.- (HIV Protease); 0 (Antiviral Agents); 0 (AG 1343); 0 (

HIV Protease Inhibitors); 0

(Isoquinolines); 0 (Ritonavir); 0 (Sulfonamides); 0 (Sulfonic

Acids); 0 (VX 478)

L30 ANSWER 11 OF 30 MEDLINE

ACCESSION NUMBER: 97112989 MEDLINE

TITLE: Mutational anatomy of an HIV-1 protease variant

conferring cross-resistance to protease inhibitors in clinical trials. Compensatory modulations of binding

and activity.

AUTHOR: Schock H B; Garsky V M; Kuo L C

CORPORATE SOURCE: Department of Antiviral Research, Merck Research

Laboratories, West Point, Pennsylvania 19486, USA.

SOURCE: JOURNAL OF BIOLOGICAL CHEMISTRY, (1996 Dec 13) 271

(50) 31957-63.

Journal code: HIV. ISSN: 0021-9258.

PUB. COUNTRY: United States

Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

FILE SEGMENT: Priority Journals; Cancer Journals

ENTRY MONTH: 9703

ENTRY WEEK: 19970304

Site-specific substitutions of as few as four amino acids (M46I/L63P/V82T/I84V) of the human immunodeficiency virus type 1 (HIV-1) protease engenders cross-resistance to a panel of protease inhibitors that are either in clinical trials or have recently been approved for HIV therapy (Condra, J. H., Schleif, W. A., Blahy, O. M., Gadryelski, L. J., Graham, D. J., Quintero, J. C., Rhodes, A., Robbins, H. L., Roth, E., Shivaprakash, M., Titus, D., Yang, T., Teppler, H., Squires, K. E., Deutsch, P. J., and Emini, E. A. (1995) Nature 374, 569-571). These four substitutions are among the prominent mutations found in primary HIV isolates obtained from patients undergoing therapy with several protease inhibitors. Two of these mutations (V82T/I84V) are located in, while the other two (M46I/L63P) are away from, the binding cleft of the enzyme. The functional role of these mutations has now been delineated in terms of their influence on the binding affinity and catalytic efficiency of the protease. We have found that the double substitutions of M46I and L63P do not affect binding but instead endow the enzyme with a catalytic efficiency significantly exceeding (110-360%) that of the wild-type enzyme. In contrast, the double substitutions of V82T and I84V are detrimental to the ability of the protease to bind and, thereby, to catalyze. When combined, the four amino acid replacements institute in the protease resistance against inhibitors and a significantly higher catalytic activity than one containing only mutations in its active site. The results suggest that in raising drug resistance, these four site-specific mutations of the protease are compensatory in function; those in the active site diminish equilibrium binding (by increasing Ki), and those away from the active site enhance catalysis (by increasing kcat/KM). This

conclusion is further supported by energy estimates in that the Gibbs free energies of binding and catalysis for the quadruple mutant are quantitatively dictated by those of the double mutants. CTCheck Tags: Human Clinical Trials Fusion Proteins, gag-pol: ME, metabolism Hydrolysis HIV Protease: CH, chemistry *HIV Protease: GE, genetics *HIV Protease Inhibitors: PD, pharmacology Indinavir: PD, pharmacology Kinetics Mutagenesis Ritonavir: PD, pharmacology Sulfonamides: PD, pharmacology CN EC 3.4.23.- (HIV Protease); 0 (Fusion Proteins, gag-pol); 0 (HIV Protease Inhibitors); 0 (Ritonavir); 0 (Sulfonamides); 0 (VX 478) 130 ANSWER 12 OF 30 MEDLINE ACCESSION NUMBER: 97046545 MEDLINE TITLE: Ritonavir. AUTHOR: Lea A P; Faulds D CORPORATE SOURCE: Adis International Limited, Auckland, New Zealand. SOURCE: DRUGS, (1996 Oct) 52 (4) 541-6; discussion 547-8. Ref: 37 Journal code: EC2. ISSN: 0012-6667. PUB. COUNTRY: New Zealand Journal; Article; (JOURNAL ARTICLE) General Review; (REVIEW) (REVIEW, TUTORIAL) LANGUAGE: English FILE SEGMENT: Priority Journals ENTRY MONTH: 9704 ENTRY WEEK: 19970401 Ritonavir is a protease inhibitor with an HIV-1 resistance profile similar to that of indinavir, but different from that of saquinavir. Ritonavir has good oral bioavailability, and may increase the bioavailability of other protease inhibitors including saquinavir, nelfinavir, indinavir and VX-478. Clinically significant drug interactions have been predicted between ritonavir and a range of medications. In patients with HIV-1 infection, ritonavir markedly reduced viral load within 2 weeks of treatment onset and also increased CD4+ cell counts. In a large placebo-controlled trial in patients with advanced HIV infection, the addition of ritonavir to existing therapy reduced the risk of mortality by 43% and clinical progression by 56% after 6.1 months. Triple therapy with ritonavir plus zidovudine, in combination with lamivudine or zalcitabine, reduced HIV viraemia to below detectable levels in most patients with acute, and some patients with advanced HIV infection in 2 small trials. Early results suggest combination therapy with ritonavir and saquinavir increases CD4+ cell counts and

AB Ritonavir is a protease inhibitor with an HIV-1 resistance profile similar to that of indinavir, but different from that of saquinavir. Ritonavir has good oral bioavailability, and may increase the bioavailability of other protease inhibitors including saquinavir,

infection.

decreases HIV RNA levels in patients with previously untreated HIV

4-14-97

=> s ritonavir/cn

L2 1 RITONAVIR/CN

=> d 12

L2 ANSWER 1 OF 1 REGISTRY COPYRIGHT 1997 ACS

RN 155213-67-5 REGISTRY

CN 2,4,7,12-Tetraazatridecan-13-oic acid, 10-hydroxy-2-methyl-5-(1-methylethyl)-1-[2-(1-methylethyl)-4-thiazolyl]-3,6-dioxo-8,11-bis(phenylmethyl)-, 5-thiazolylmethyl ester, [5S-(5R*,8R*,10R*,11R*)]- (9CI) (CA INDEX NAME)

OTHER NAMES:

CN A 84538

CN Abbott 84538

CN ABT 538

CN Norvir

CN Ritonavir

FS STEREOSEARCH

MF C37 H48 N6 O5 S2

CI COM

SR CAS Registry Services

LC STN Files: BIOBUSINESS, BIOSIS, CA, CAPLUS, CEN, CHEMLIST, CIN, DDFU, DRUGNL, DRUGPAT, DRUGU, DRUGUPDATES, EMBASE, IPA, PHAR, PROMT, TOXLINE, TOXLIT, USAN, USPATFULL

Absolute stereochemistry.

35 REFERENCES IN FILE CA (1967 TO DATE)
35 REFERENCES IN FILE CAPLUS (1967 TO DATE)

=> s 13 and cyclosporine 4007 CYCLOSPORINE

1 L3 AND CYCLOSPORINE L5

=> d bib ab

L5 ANSWER 1 OF 1 CAPLUS COPYRIGHT 1997 ACS

1997:184660 CAPLUS ΑN

DN 126:166463

Use of ritonavir (ABT-538) for improving the pharmacokinetics of TΙ drugs metabolized by cytochrome P450 in a method of treating aids

Norbeck, Daniel W.; Kempf, Dale J.; Leonard, John M.; Bertz, Richard ΙN J.

Abbott Laboratories, USA PΑ

PCT Int. Appl., 28 pp. SO

CODEN: PIXXD2

WO 9701349 A1 970116 PΙ

DS W: AU, CA, IS, JP, KR, MX

RW: AT, BE, CH, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT,

ΑI WO 96-US11015 960628

PRAI US 95-654 950629

US 95-3849 950915

DT Patent

LΑ English

A method is disclosed for improving the pharmacokinetics of a drug AΒ which is metabolized by cytochrome P 450 monooxygenase by use of ritonavir. HIV inhibitory action is also claimed by combinations of ritonavir with protease inhibitors whose pharmacokinetics are modulated by ritanovir via its inhibitory action on cytochrome P 450.

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=> s wo9414436/pn
             1 W09414436/PN
                 (WO9414436/PN)
=> d ibib ab
     ANSWER 1 OF 1 WPIDS
                            COPYRIGHT 1997 DERWENT INFORMATION LTD
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                      94-234319 [28]
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ACCESSION NUMBER:
CROSS REFERENCE:
                      90-377452 [51];
                                        92-176578 [22];
                                                         92-348620 [42];
                      93-350874 [44];
                                        93-386428 [48];
                                                         94-357440 [44]
DOC. NO. CPI:
                      C94-106528
                      New di heterocyclyl-substd. carbonate cpds. - used
TITLE:
                      as HIV protease inhibiting anti-retroviral agents
                      esp. for treating AIDS.
DERWENT CLASS:
                      B03
                      KEMPF, D J; NORBECK, D W; SHAM, H L; ZHAO, C;
INVENTOR(S):
                      COOPER, A J; HAIGHT, A R; RENO, D S; SOWIN, T J;
                      ALLEN, M S; COPPER, A J; TIEN, J J
                      (ABBO) ABBOTT LAB
PATENT ASSIGNEE(S):
COUNTRY COUNT:
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PATENT INFORMATION:
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APPLICATION DETAILS:

WO	9414436	A1				WO	93-US12326	931216
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FILING DETAILS:

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29; US PRIOR 890908; US 89-355945 890523; US 89-405604 891222; US 90-518730 900509; US 89-456124 901120; US 91-746020 90-616170 910815; US 911023; US 95-410996 950327; US 91-777626 950425; US 95-411032 950327; US 95-423387 95-417304 950405; US 95-412253 950328; US 95-412821 950329; US 95-413290 950330; US 950404; US 95-412438 950329; US 95-416272 950404; US 95-416259 950404 95-416607

AB WO 9414436 A UPAB: 960918

O-(Heterocyclyl-alkyl) - N-(Heterocyclyl-substd. carbonylamino-alkyl) carbamates of formula (I) and their salts, esters and prodrugs are new. R1 = thiazolyl, oxazolyl, isoxazolyl or isothiazolyl, all mono-substd. by Q; Q = lower alkyl, lower alkenyl, cycloalkyl, cycloalkylalkyl, cycloalkenyl, cycloalkenylalkyl, Het, Het-alkyl, alkoxyalkyl, thioalkoxyalkyl, alkylamino, dialkylamino, phenyl (opt. substd. by halo, lower alkyl, OH, alkoxy or thioalkoxy), phenyl alkyl (opt. ring-substd. as for phenyl), dialkylaminoalkyl, alkoxy or thioalkoxy; Het = aziridinyl, azetidinyl, pyrrolidinyl, piperidinyl, piperazinyl, morpholinyl, thiomorpholinyl, thiazolyl, oxazolyl, isoxazolyl, isothiazolyl, pyridinyl, pyrimidinyl, pyridazinyl or pyrazinyl (all opt. substd. by halo, lower alkyl, OH, alkoxy or thioalkoxy); n = 1-3; R2 = H or lower alkyl; R3 = loweralkyl; R4, R4a = phenyl, thiazolyl or oxazolyl (all opt. substd. by halo, lower alkyl, OH, alkoxy or thioalkoxy); R6 = H or lower alkoxy; R7 = chiazolyl, oxazolyl, isoxazolyl, or isothiazolyl, (all opt. substd. by lower alkyl); one of X, Y = H and the other = OH; or X = Y = OH Z = direct bond, O, S, CH2 or NR8; R8 = lower alkyl,cycloalkyl, OH or NHR8a; R8a = H, lower alkyl or N-protecting gp.; provided that, in cpds. where only one of X and Y = OH, X = H and Y = OH when R7 is unsubstd. and either Z = NR8 or R3 = Me lower = up to 6C.

USE/ADVANTAGE - (I) are retroviral protease inhibitors, esp. HIV protease inhibitor (claimed), effective in vitro or in vivo. (I) inhibit retroviruses, esp. HIV, in vivo and are useful for treatment or prophylaxis of diseases caused by retroviruses, esp. AIDS or HIV

infection. (I) may be ree of the adverse effects of cover anti-AIDS agents, e.g. low platest count, renal toxicity and be marrow cytopenia.

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